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(54) Title: COMPOSITIONS AND METHODS FOR THERAPY AND DIAGNOSIS OF BREAST CANCER

(57) Abstract: Compositions and methods for the therapy and diagnosis of cancer, such as breast cancer, are disclosed. Compositions may comprise one or more breast tumor proteins, immunogenic portions thereof, or polynucleotides that encode such portions. Alternatively, a therapeutic composition may comprise an antigen presenting cell that expresses a breast tumor protein, or a T cell that is specific for cells expressing such a protein. Such compositions may be used, for example, for the prevention and treatment of diseases such as breast cancer. Diagnostic methods based on detecting a breast tumor protein, or mRNA encoding such a protein, in a sample are also provided.

**COMPOSITIONS AND METHODS FOR THERAPY AND DIAGNOSIS
OF BREAST CANCER**

TECHNICAL FIELD

The present invention relates generally to therapy and diagnosis of cancer, such as breast cancer. The invention is more specifically related to polypeptides comprising at least a portion of a breast tumor protein, and to polynucleotides encoding such polypeptides. Such polypeptides and polynucleotides may be used in vaccines and pharmaceutical compositions for prevention and treatment of breast cancer, and for the diagnosis and monitoring of such cancers.

10 BACKGROUND OF THE INVENTION

Breast cancer is a significant health problem for women in the United States and throughout the world. Although advances have been made in detection and treatment of the disease, breast cancer remains the second leading cause of cancer-related deaths in women, affecting more than 180,000 women in the United States each year. For women in North America, the life-time odds of getting breast cancer are now one in eight.

No vaccine or other universally successful method for the prevention or treatment of breast cancer is currently available. Management of the disease currently relies on a combination of early diagnosis (through routine breast screening procedures) and aggressive treatment, which may include one or more of a variety of treatments such as surgery, radiotherapy, chemotherapy and hormone therapy. The course of treatment for a particular breast cancer is often selected based on a variety of prognostic parameters, including an analysis of specific tumor markers. See, e.g., Porter-Jordan and Lippman, *Breast Cancer* 8:73-100 (1994). However, the use of established markers often leads to a result that is difficult to interpret, and the high mortality observed in breast cancer patients indicates that improvements are needed in the treatment, diagnosis and prevention of the disease.

Accordingly, there is a need in the art for improved methods for therapy and diagnosis of breast cancer. The present invention fulfills these needs and further provides other related advantages.

SUMMARY OF THE INVENTION

5 Briefly stated, the present invention provides compositions and methods for the diagnosis and therapy of cancer, such as breast cancer. In one aspect, the present invention provides polypeptides comprising at least a portion of a breast tumor protein, or a variant thereof. Certain portions and other variants are immunogenic, such that the ability of the variant to react with antigen-specific antisera is not substantially 10 diminished. Within certain embodiments, the polypeptide comprises a sequence that is encoded by a polynucleotide sequence selected from the group consisting of: (a) sequences recited in SEQ ID NO: 1-38, 42-204, 205, 207 and 210-290; (b) variants of a sequence recited in SEQ ID NO: 1-38, 42-204, 205, 207 and 210-290; and (c) complements of a sequence of (a) or (b).

15 The present invention further provides polynucleotides that encode a polypeptide as described above, or a portion thereof (such as a portion encoding at least 15 amino acid residues of a breast tumor protein), expression vectors comprising such polynucleotides and host cells transformed or transfected with such expression vectors.

20 Within other aspects, the present invention provides pharmaceutical compositions comprising a polypeptide or polynucleotide as described above and a physiologically acceptable carrier.

Within a related aspect of the present invention, vaccines for prophylactic or therapeutic use are provided. Such vaccines comprise a polypeptide or polynucleotide as described above and an immunostimulant.

25 The present invention further provides pharmaceutical compositions that comprise: (a) an antibody or antigen-binding fragment thereof that specifically binds to a breast tumor protein; and (b) a physiologically acceptable carrier.

Within further aspects, the present invention provides pharmaceutical compositions comprising: (a) an antigen presenting cell that expresses a polypeptide as 30 described above and (b) a pharmaceutically acceptable carrier or excipient. Antigen

presenting cells include dendritic cells, macrophages, monocytes, fibroblasts and B cells.

Within related aspects, vaccines are provided that comprise: (a) an antigen presenting cell that expresses a polypeptide as described above and (b) an
5 immunostimulant.

The present invention further provides, in other aspects, fusion proteins that comprise at least one polypeptide as described above, as well as polynucleotides encoding such fusion proteins.

Within related aspects, pharmaceutical compositions comprising a fusion
10 protein, or a polynucleotide encoding a fusion protein, in combination with a physiologically acceptable carrier are provided.

Vaccines are further provided, within other aspects, that comprise a fusion protein, or a polynucleotide encoding a fusion protein, in combination with an immunostimulant.

15 Within further aspects, the present invention provides methods for inhibiting the development of a cancer in a patient, comprising administering to a patient a pharmaceutical composition or vaccine as recited above.

The present invention further provides, within other aspects, methods for removing tumor cells from a biological sample, comprising contacting a biological
20 sample with T cells that specifically react with a breast tumor protein, wherein the step of contacting is performed under conditions and for a time sufficient to permit the removal of cells expressing the protein from the sample.

Within related aspects, methods are provided for inhibiting the development of a cancer in a patient, comprising administering to a patient a biological
25 sample treated as described above.

Methods are further provided, within other aspects, for stimulating and/or expanding T cells specific for a breast tumor protein, comprising contacting T cells with one or more of: (i) a polypeptide as described above; (ii) a polynucleotide encoding such a polypeptide; and/or (iii) an antigen presenting cell that expresses such a
30 polypeptide; under conditions and for a time sufficient to permit the stimulation and/or

expansion of T cells. Isolated T cell populations comprising T cells prepared as described above are also provided.

Within further aspects, the present invention provides methods for inhibiting the development of a cancer in a patient, comprising administering to a 5 patient an effective amount of a T cell population as described above.

The present invention further provides methods for inhibiting the development of a cancer in a patient, comprising the steps of: (a) incubating CD4⁺ and/or CD8⁺ T cells isolated from a patient with one or more of: (i) a polypeptide comprising at least an immunogenic portion of a breast tumor protein; (ii) a 10 polynucleotide encoding such a polypeptide; and (iii) an antigen-presenting cell that expresses such a polypeptide; and (b) administering to the patient an effective amount of the proliferated T cells, and thereby inhibiting the development of a cancer in the patient. Proliferated cells may, but need not, be cloned prior to administration to the patient.

15 Within further aspects, the present invention provides methods for determining the presence or absence of a cancer in a patient, comprising: (a) contacting a biological sample obtained from a patient with a binding agent that binds to a polypeptide as recited above; (b) detecting in the sample an amount of polypeptide that binds to the binding agent; and (c) comparing the amount of polypeptide with a 20 predetermined cut-off value, and therefrom determining the presence or absence of a cancer in the patient. Within preferred embodiments, the binding agent is an antibody, more preferably a monoclonal antibody. The cancer may be breast cancer.

The present invention also provides, within other aspects, methods for monitoring the progression of a cancer in a patient. Such methods comprise the steps 25 of: (a) contacting a biological sample obtained from a patient at a first point in time with a binding agent that binds to a polypeptide as recited above; (b) detecting in the sample an amount of polypeptide that binds to the binding agent; (c) repeating steps (a) and (b) using a biological sample obtained from the patient at a subsequent point in time; and (d) comparing the amount of polypeptide detected in step (c) with the amount 30 detected in step (b) and therefrom monitoring the progression of the cancer in the patient.

The present invention further provides, within other aspects, methods for determining the presence or absence of a cancer in a patient, comprising the steps of: (a) contacting a biological sample obtained from a patient with an oligonucleotide that hybridizes to a polynucleotide that encodes a breast tumor protein; (b) detecting in the sample a level of a polynucleotide, preferably mRNA, that hybridizes to the oligonucleotide; and (c) comparing the level of polynucleotide that hybridizes to the oligonucleotide with a predetermined cut-off value, and therefrom determining the presence or absence of a cancer in the patient. Within certain embodiments, the amount of mRNA is detected via polymerase chain reaction using, for example, at least one oligonucleotide primer that hybridizes to a polynucleotide encoding a polypeptide as recited above, or a complement of such a polynucleotide. Within other embodiments, the amount of mRNA is detected using a hybridization technique, employing an oligonucleotide probe that hybridizes to a polynucleotide that encodes a polypeptide as recited above, or a complement of such a polynucleotide.

In related aspects, methods are provided for monitoring the progression of a cancer in a patient, comprising the steps of: (a) contacting a biological sample obtained from a patient with an oligonucleotide that hybridizes to a polynucleotide that encodes a breast tumor protein; (b) detecting in the sample an amount of a polynucleotide that hybridizes to the oligonucleotide; (c) repeating steps (a) and (b) using a biological sample obtained from the patient at a subsequent point in time; and (d) comparing the amount of polynucleotide detected in step (c) with the amount detected in step (b) and therefrom monitoring the progression of the cancer in the patient.

Within further aspects, the present invention provides antibodies, such as monoclonal antibodies, that bind to a polypeptide as described above, as well as diagnostic kits comprising such antibodies. Diagnostic kits comprising one or more oligonucleotide probes or primers as described above are also provided.

These and other aspects of the present invention will become apparent upon reference to the following detailed description. All references disclosed herein are hereby incorporated by reference in their entirety as if each was incorporated individually.

SEQUENCE IDENTIFIERS

SEQ ID NO: 1 is the determined cDNA sequence for clone 26915.

SEQ ID NO: 2 is the determined cDNA sequence for clone 26914.

SEQ ID NO: 3 is the determined cDNA sequence for clone 26673.

5 SEQ ID NO: 4 is the determined cDNA sequence for clone 26672.

SEQ ID NO: 5 is the determined cDNA sequence for clone 26671.

SEQ ID NO: 6 is the determined cDNA sequence for clone 26670.

SEQ ID NO: 7 is the determined cDNA sequence for clone 26669.

SEQ ID NO: 8 is a first determined cDNA sequence for clone 26668.

10 SEQ ID NO: 9 is a second determined cDNA sequence for clone 26668.

SEQ ID NO: 10 is the determined cDNA sequence for clone 26667.

SEQ ID NO: 11 is the determined cDNA sequence for clone 26666.

SEQ ID NO: 12 is the determined cDNA sequence for clone 26665.

SEQ ID NO: 13 is the determined cDNA sequence for clone 26664.

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SEQ ID NO: 15 is the determined cDNA sequence for clone 26661.

SEQ ID NO: 16 is the determined cDNA sequence for clone 26660.

SEQ ID NO: 17 is the determined cDNA sequence for clone 26603.

SEQ ID NO: 18 is the determined cDNA sequence for clone 26601.

20 SEQ ID NO: 19 is the determined cDNA sequence for clone 26600.

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SEQ ID NO: 21 is the determined cDNA sequence for clone 26586.

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25 SEQ ID NO: 24 is the determined cDNA sequence for clone 26580.

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30 SEQ ID NO: 29 is the determined cDNA sequence for clone 26573.

SEQ ID NO: 30 is the determined cDNA sequence for clone 25612.

SEQ ID NO: 31 is the determined cDNA sequence for clone 22295.

SEQ ID NO: 32 is the determined cDNA sequence for clone 22301.

SEQ ID NO: 33 is the determined cDNA sequence for clone 22298.

SEQ ID NO: 34 is the determined cDNA sequence for clone 22297.

5 SEQ ID NO: 35 is the determined cDNA sequence for clone 22303.

SEQ ID NO: 36 is the determined cDNA sequence for a first GABA_A receptor clone.

SEQ ID NO: 37 is the determined cDNA sequence for a second GABA_A receptor clone.

10 SEQ ID NO: 38 is the determined cDNA sequence for a third GABA_A receptor clone.

SEQ ID NO: 39 is the amino acid sequence encoded by SEQ ID NO: 36.

SEQ ID NO: 40 is the amino acid sequence encoded by SEQ ID NO: 37.

SEQ ID NO: 41 is the amino acid sequence encoded by SEQ ID NO: 38.

15 SEQ ID NO: 42 is the determined cDNA sequence for contig 1.

SEQ ID NO: 43 is the determined cDNA sequence for contig 2.

SEQ ID NO: 44 is the determined cDNA sequence for contig 3.

SEQ ID NO: 45 is the determined cDNA sequence for contig 4.

SEQ ID NO: 46 is the determined cDNA sequence for contig 5.

20 SEQ ID NO: 47 is the determined cDNA sequence for contig 6.

SEQ ID NO: 48 is the determined cDNA sequence for contig 7.

SEQ ID NO: 49 is the determined cDNA sequence for contig 8.

SEQ ID NO: 50 is the determined cDNA sequence for contig 9.

SEQ ID NO: 51 is the determined cDNA sequence for contig 10.

25 SEQ ID NO: 52 is the determined cDNA sequence for contig 11.

SEQ ID NO: 53 is the determined cDNA sequence for contig 12.

SEQ ID NO: 54 is the determined cDNA sequence for contig 13.

SEQ ID NO: 55 is the determined cDNA sequence for contig 14.

SEQ ID NO: 56 is the determined cDNA sequence for contig 15.

30 SEQ ID NO: 57 is the determined cDNA sequence for contig 16.

SEQ ID NO: 58 is the determined cDNA sequence for contig 17.

SEQ ID NO: 59 is the determined cDNA sequence for contig 18.
SEQ ID NO: 60 is the determined cDNA sequence for contig 19.
SEQ ID NO: 61 is the determined cDNA sequence for contig 20.
SEQ ID NO: 62 is the determined cDNA sequence for contig 21.
5 SEQ ID NO: 63 is the determined cDNA sequence for contig 22.
SEQ ID NO: 64 is the determined cDNA sequence for contig 23.
SEQ ID NO: 65 is the determined cDNA sequence for contig 24.
SEQ ID NO: 66 is the determined cDNA sequence for contig 25.
SEQ ID NO: 67 is the determined cDNA sequence for contig 26.
10 SEQ ID NO: 68 is the determined cDNA sequence for contig 27.
SEQ ID NO: 69 is the determined cDNA sequence for contig 28.
SEQ ID NO: 70 is the determined cDNA sequence for contig 29.
SEQ ID NO: 71 is the determined cDNA sequence for contig 30.
SEQ ID NO: 72 is the determined cDNA sequence for contig 31.
15 SEQ ID NO: 73 is the determined cDNA sequence for contig 32.
SEQ ID NO: 74 is the determined cDNA sequence for contig 33.
SEQ ID NO: 75 is the determined cDNA sequence for contig 34.
SEQ ID NO: 76 is the determined cDNA sequence for contig 35.
SEQ ID NO: 77 is the determined cDNA sequence for contig 36.
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SEQ ID NO: 82 is the determined cDNA sequence for contig 41.
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SEQ ID NO: 84 is the determined cDNA sequence for contig 43.
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SEQ ID NO: 85 is the determined cDNA sequence for contig 45.
SEQ ID NO: 85 is the determined cDNA sequence for contig 46.
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SEQ ID NO: 89 is the determined cDNA sequence for contig 48.

SEQ ID NO: 90 is the determined cDNA sequence for contig 49.
SEQ ID NO: 91 is the determined cDNA sequence for contig 50.
SEQ ID NO: 92 is the determined cDNA sequence for contig 51.
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SEQ ID NO: 97 is the determined cDNA sequence for contig 56.
SEQ ID NO: 98 is the determined cDNA sequence for contig 57.
10 SEQ ID NO: 99 is the determined cDNA sequence for contig 58.
SEQ ID NO: 100 is the determined cDNA sequence for contig 59.
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SEQ ID NO: 102 is the determined cDNA sequence for contig 61.
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SEQ ID NO: 107 is the determined cDNA sequence for contig 66.
SEQ ID NO: 108 is the determined cDNA sequence for contig 67.
20 SEQ ID NO: 109 is the determined cDNA sequence for contig 68.
SEQ ID NO: 110 is the determined cDNA sequence for contig 69.
SEQ ID NO: 111 is the determined cDNA sequence for contig 70.
SEQ ID NO: 112 is the determined cDNA sequence for contig 71.
SEQ ID NO: 113 is the determined cDNA sequence for contig 72.
25 SEQ ID NO: 114 is the determined cDNA sequence for contig 73.
SEQ ID NO: 115 is the determined cDNA sequence for contig 74.
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SEQ ID NO: 117 is the determined cDNA sequence for contig 76.
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30 SEQ ID NO: 119 is the determined cDNA sequence for contig 78.
SEQ ID NO: 120 is the determined cDNA sequence for contig 79.

SEQ ID NO: 121 is the determined cDNA sequence for contig 80.
SEQ ID NO: 122 is the determined cDNA sequence for contig 81.
SEQ ID NO: 123 is the determined cDNA sequence for contig 82.
SEQ ID NO: 124 is the determined cDNA sequence for contig 83.
5 SEQ ID NO: 125 is the determined cDNA sequence for contig 84.
SEQ ID NO: 126 is the determined cDNA sequence for contig 85.
SEQ ID NO: 127 is the determined cDNA sequence for contig 86.
SEQ ID NO: 128 is the determined cDNA sequence for contig 87.
SEQ ID NO: 129 is the determined cDNA sequence for contig 88.
10 SEQ ID NO: 130 is the determined cDNA sequence for contig 89.
SEQ ID NO: 131 is the determined cDNA sequence for contig 90.
SEQ ID NO: 132 is the determined cDNA sequence for contig 91.
SEQ ID NO: 133 is the determined cDNA sequence for contig 92.
SEQ ID NO: 134 is the determined cDNA sequence for contig 93.
15 SEQ ID NO: 135 is the determined cDNA sequence for contig 94.
SEQ ID NO: 136 is the determined cDNA sequence for contig 95.
SEQ ID NO: 137 is the determined cDNA sequence for contig 96.
SEQ ID NO: 138 is the determined cDNA sequence for clone 47589.
SEQ ID NO: 139 is the determined cDNA sequence for clone 47578.
20 SEQ ID NO: 140 is the determined cDNA sequence for clone 47602.
SEQ ID NO: 141 is the determined cDNA sequence for clone 47593.
SEQ ID NO: 142 is the determined cDNA sequence for clone 47583.
SEQ ID NO: 143 is the determined cDNA sequence for clone 47624.
SEQ ID NO: 144 is the determined cDNA sequence for clone 47622.
25 SEQ ID NO: 145 is the determined cDNA sequence for clone 47649.
SEQ ID NO: 146 is the determined cDNA sequence for clone 48955.
SEQ ID NO: 147 is the determined cDNA sequence for clone 48962.
SEQ ID NO: 148 is the determined cDNA sequence for clone 48964.
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SEQ ID NO: 154 is the determined cDNA sequence for clone 48931.
SEQ ID NO: 155 is the determined cDNA sequence for clone 48935.
5 SEQ ID NO: 156 is the determined cDNA sequence for clone 48940.
SEQ ID NO: 157 is the determined cDNA sequence for clone 48936.
SEQ ID NO: 158 is the determined cDNA sequence for clone 48930.
SEQ ID NO: 159 is the determined cDNA sequence for clone 48956.
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SEQ ID NO: 164 is the determined cDNA sequence for clone 48984.
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15 SEQ ID NO: 166 is the determined cDNA sequence for clone 48978.
SEQ ID NO: 167 is the determined cDNA sequence for clone 48968.
SEQ ID NO: 168 is the determined cDNA sequence for clone 48929.
SEQ ID NO: 169 is the determined cDNA sequence for clone 48937.
SEQ ID NO: 170 is the determined cDNA sequence for clone 48982.
20 SEQ ID NO: 171 is the determined cDNA sequence for clone 48983.
SEQ ID NO: 172 is the determined cDNA sequence for clone 48997.
SEQ ID NO: 173 is the determined cDNA sequence for clone 48992.
SEQ ID NO: 174 is the determined cDNA sequence for clone 49006.
SEQ ID NO: 175 is the determined cDNA sequence for clone 48994.
25 SEQ ID NO: 176 is the determined cDNA sequence for clone 49013.
SEQ ID NO: 177 is the determined cDNA sequence for clone 49008.
SEQ ID NO: 178 is the determined cDNA sequence for clone 48990.
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SEQ ID NO: 180 is the determined cDNA sequence for clone 49014.
30 SEQ ID NO: 181 is the determined cDNA sequence for clone 48988.
SEQ ID NO: 182 is the determined cDNA sequence for clone 49018.

SEQ ID NO: 183 is the determined cDNA sequence for clone 6921.

SEQ ID NO: 184 is the determined cDNA sequence for clone 6837.

SEQ ID NO: 185 is the determined cDNA sequence for clone 6840.

SEQ ID NO: 186 is the determined cDNA sequence for clone 6844.

5 SEQ ID NO: 187 is the determined cDNA sequence for clone 6854.

SEQ ID NO: 188 is the determined cDNA sequence for clone 6872.

SEQ ID NO: 189 is the determined cDNA sequence for clone 6906.

SEQ ID NO: 190 is the determined cDNA sequence for clone 6908.

SEQ ID NO: 191 is the determined cDNA sequence for clone 6910.

10 SEQ ID NO: 192 is the determined cDNA sequence for clone 6912.

SEQ ID NO: 193 is the determined cDNA sequence for clone 6913.

SEQ ID NO: 194 is the determined cDNA sequence for clone 6914.

SEQ ID NO: 195 is the determined cDNA sequence for clone 6916.

SEQ ID NO: 196 is the determined cDNA sequence for clone 6918.

15 SEQ ID NO: 197 is the determined cDNA sequence for clone 6924.

SEQ ID NO: 198 is the determined cDNA sequence for clone 6928.

SEQ ID NO: 199 is the determined cDNA sequence for clone 6978A.

SEQ ID NO: 200 is the determined cDNA sequence for clone 6978B.

SEQ ID NO: 201 is the determined cDNA sequence for clone 6982A.

20 SEQ ID NO: 202 is the determined cDNA sequence for clone 6982B.

SEQ ID NO: 203 is the determined cDNA sequence for clone 6850.

SEQ ID NO: 204 is the determined cDNA sequence for clone 6860.

SEQ ID NO: 205 is the determined cDNA sequence for O772P.

SEQ ID NO: 206 is the amino acid sequence encoded by SEQ ID NO:

25 205.

SEQ ID NO: 207 is the full-length cDNA sequence for O8E.

SEQ ID NO: 208 is a first amino acid sequence encoded by SEQ ID NO:

207.

SEQ ID NO: 209 is a second amino acid sequence encoded by SEQ ID

30 NO: 209.

SEQ ID NO: 210-290 are determined cDNA sequence of breast-tumor specific clones.

DETAILED DESCRIPTION OF THE INVENTION

As noted above, the present invention is generally directed to compositions and methods for using the compositions, for example in the therapy and diagnosis of cancer, such as breast cancer. Certain illustrative compositions described herein include breast tumor polypeptides, polynucleotides encoding such polypeptides, binding agents such as antibodies, antigen presenting cells (APCs) and/or immune system cells (*e.g.*, T cells). A "breast tumor protein," as the term is used herein, refers generally to a protein that is expressed in breast tumor cells at a level that is at least two fold, and preferably at least five fold, greater than the level of expression in other normal tissues, as determined using a representative assay provided herein. Certain breast tumor proteins are tumor proteins that react detectably (within an immunoassay, such as an ELISA or Western blot) with antisera of a patient afflicted with breast cancer.

Therefore, in accordance with the above, and as described further below, the present invention provides illustrative polynucleotide compositions having sequences set forth in SEQ ID NO:1-38, 42-204, 205, 207 and 210-290, polypeptides encoded by such polynucleotides, antibody compositions capable of binding such polypeptides, and numerous additional embodiments employing such compositions, for example in the detection, diagnosis and/or therapy of human breast cancer.

POLYNUCLEOTIDE COMPOSITIONS

As used herein, the terms "DNA segment" and "polynucleotide" refer to a DNA molecule that has been isolated free of total genomic DNA of a particular species. Therefore, a DNA segment encoding a polypeptide refers to a DNA segment that contains one or more coding sequences yet is substantially isolated away from, or purified free from, total genomic DNA of the species from which the DNA segment is obtained. Included within the terms "DNA segment" and "polynucleotide" are DNA

segments and smaller fragments of such segments, and also recombinant vectors, including, for example, plasmids, cosmids, phagemids, phage, viruses, and the like.

As will be understood by those skilled in the art, the DNA segments of this invention can include genomic sequences, extra-genomic and plasmid-encoded 5 sequences and smaller engineered gene segments that express, or may be adapted to express, proteins, polypeptides, peptides and the like. Such segments may be naturally isolated, or modified synthetically by the hand of man.

"Isolated," as used herein, means that a polynucleotide is substantially away from other coding sequences, and that the DNA segment does not contain large 10 portions of unrelated coding DNA, such as large chromosomal fragments or other functional genes or polypeptide coding regions. Of course, this refers to the DNA segment as originally isolated, and does not exclude genes or coding regions later added to the segment by the hand of man.

As will be recognized by the skilled artisan, polynucleotides may be 15 single-stranded (coding or antisense) or double-stranded, and may be DNA (genomic, cDNA or synthetic) or RNA molecules. RNA molecules include HnRNA molecules, which contain introns and correspond to a DNA molecule in a one-to-one manner, and mRNA molecules, which do not contain introns. Additional coding or non-coding sequences may, but need not, be present within a polynucleotide of the present 20 invention, and a polynucleotide may, but need not, be linked to other molecules and/or support materials.

Polynucleotides may comprise a native sequence (*i.e.*, an endogenous sequence that encodes a breast tumor protein or a portion thereof) or may comprise a variant, or a biological or antigenic functional equivalent of such a sequence. 25 Polynucleotide variants may contain one or more substitutions, additions, deletions and/or insertions, as further described below, preferably such that the immunogenicity of the encoded polypeptide is not diminished, relative to a native tumor protein. The effect on the immunogenicity of the encoded polypeptide may generally be assessed as described herein. The term "variants" also encompasses homologous genes of 30 xenogenic origin.

When comparing polynucleotide or polypeptide sequences, two sequences are said to be "identical" if the sequence of nucleotides or amino acids in the two sequences is the same when aligned for maximum correspondence, as described below. Comparisons between two sequences are typically performed by comparing the 5 sequences over a comparison window to identify and compare local regions of sequence similarity. A "comparison window" as used herein, refers to a segment of at least about 20 contiguous positions, usually 30 to about 75, 40 to about 50, in which a sequence may be compared to a reference sequence of the same number of contiguous positions after the two sequences are optimally aligned.

10 Optimal alignment of sequences for comparison may be conducted using the Megalign program in the Lasergene suite of bioinformatics software (DNASTAR, Inc., Madison, WI), using default parameters. This program embodies several alignment schemes described in the following references: Dayhoff, M.O. (1978) A model of evolutionary change in proteins – Matrices for detecting distant relationships.

15 In Dayhoff, M.O. (ed.) *Atlas of Protein Sequence and Structure*, National Biomedical Research Foundation, Washington DC Vol. 5, Suppl. 3, pp. 345-358; Hein J. (1990) Unified Approach to Alignment and Phylogenies pp. 626-645 *Methods in Enzymology* vol. 183, Academic Press, Inc., San Diego, CA; Higgins, D.G. and Sharp, P.M. (1989) *CABIOS* 5:151-153; Myers, E.W. and Muller W. (1988) *CABIOS* 4:11-17; Robinson,

20 E.D. (1971) *Comb. Theor* 11:105; Santou, N. Nes, M. (1987) *Mol. Biol. Evol.* 4:406-425; Sneath, P.H.A. and Sokal, R.R. (1973) *Numerical Taxonomy – the Principles and Practice of Numerical Taxonomy*, Freeman Press, San Francisco, CA; Wilbur, W.J. and Lipman, D.J. (1983) *Proc. Natl. Acad. Sci. USA* 80:726-730.

25 Alternatively, optimal alignment of sequences for comparison may be conducted by the local identity algorithm of Smith and Waterman (1981) *Add. APL. Math* 2:482, by the identity alignment algorithm of Needleman and Wunsch (1970) *J. Mol. Biol.* 48:443, by the search for similarity methods of Pearson and Lipman (1988) *Proc. Natl. Acad. Sci. USA* 85: 2444, by computerized implementations of these algorithms (GAP, BESTFIT, BLAST, FASTA, and TFASTA in the Wisconsin Genetics

30 Software Package, Genetics Computer Group (GCG), 575 Science Dr., Madison, WI), or by inspection.

One preferred example of algorithms that are suitable for determining percent sequence identity and sequence similarity are the BLAST and BLAST 2.0 algorithms, which are described in Altschul et al. (1977) *Nucl. Acids Res.* 25:3389-3402 and Altschul et al. (1990) *J. Mol. Biol.* 215:403-410, respectively. BLAST and BLAST 5 2.0 can be used, for example with the parameters described herein, to determine percent sequence identity for the polynucleotides and polypeptides of the invention. Software for performing BLAST analyses is publicly available through the National Center for Biotechnology Information. In one illustrative example, cumulative scores can be calculated using, for nucleotide sequences, the parameters M (reward score for a pair of 10 matching residues; always >0) and N (penalty score for mismatching residues; always <0). For amino acid sequences, a scoring matrix can be used to calculate the cumulative score. Extension of the word hits in each direction are halted when: the cumulative alignment score falls off by the quantity X from its maximum achieved value; the cumulative score goes to zero or below, due to the accumulation of one or 15 more negative-scoring residue alignments; or the end of either sequence is reached. The BLAST algorithm parameters W, T and X determine the sensitivity and speed of the alignment. The BLASTN program (for nucleotide sequences) uses as defaults a wordlength (W) of 11, and expectation (E) of 10, and the BLOSUM62 scoring matrix (see Henikoff and Henikoff (1989) *Proc. Natl. Acad. Sci. USA* 89:10915) alignments, 20 (B) of 50, expectation (E) of 10, M=5, N=-4 and a comparison of both strands.

Preferably, the “percentage of sequence identity” is determined by comparing two optimally aligned sequences over a window of comparison of at least 20 positions, wherein the portion of the polynucleotide or polypeptide sequence in the comparison window may comprise additions or deletions (*i.e.*, gaps) of 20 percent or 25 less, usually 5 to 15 percent, or 10 to 12 percent, as compared to the reference sequences (which does not comprise additions or deletions) for optimal alignment of the two sequences. The percentage is calculated by determining the number of positions at which the identical nucleic acid bases or amino acid residue occurs in both sequences to yield the number of matched positions, dividing the number of matched positions by the 30 total number of positions in the reference sequence (*i.e.*, the window size) and multiplying the results by 100 to yield the percentage of sequence identity.

Therefore, the present invention encompasses polynucleotide and polypeptide sequences having substantial identity to the sequences disclosed herein, for example those comprising at least 50% sequence identity, preferably at least 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 96%, 97%, 98%, or 99% or higher, sequence 5 identity compared to a polynucleotide or polypeptide sequence of this invention using the methods described herein, (e.g., BLAST analysis using standard parameters, as described below). One skilled in this art will recognize that these values can be appropriately adjusted to determine corresponding identity of proteins encoded by two nucleotide sequences by taking into account codon degeneracy, amino acid similarity, 10 reading frame positioning and the like.

In additional embodiments, the present invention provides isolated polynucleotides and polypeptides comprising various lengths of contiguous stretches of sequence identical to or complementary to one or more of the sequences disclosed herein. For example, polynucleotides are provided by this invention that comprise at 15 least about 15, 20, 30, 40, 50, 75, 100, 150, 200, 300, 400, 500 or 1000 or more contiguous nucleotides of one or more of the sequences disclosed herein as well as all intermediate lengths there between. It will be readily understood that "intermediate lengths", in this context, means any length between the quoted values, such as 16, 17, 18, 19, *etc.*; 21, 22, 23, *etc.*; 30, 31, 32, *etc.*; 50, 51, 52, 53, *etc.*; 100, 101, 102, 103, 20 *etc.*; 150, 151, 152, 153, *etc.*; including all integers through 200-500; 500-1,000, and the like.

The polynucleotides of the present invention, or fragments thereof, regardless of the length of the coding sequence itself, may be combined with other DNA sequences, such as promoters, polyadenylation signals, additional restriction 25 enzyme sites, multiple cloning sites, other coding segments, and the like, such that their overall length may vary considerably. It is therefore contemplated that a nucleic acid fragment of almost any length may be employed, with the total length preferably being limited by the ease of preparation and use in the intended recombinant DNA protocol. For example, illustrative DNA segments with total lengths of about 10,000, about 5000, 30 about 3000, about 2,000, about 1,000, about 500, about 200, about 100, about 50 base

pairs in length, and the like, (including all intermediate lengths) are contemplated to be useful in many implementations of this invention.

In other embodiments, the present invention is directed to polynucleotides that are capable of hybridizing under moderately stringent conditions to 5 a polynucleotide sequence provided herein, or a fragment thereof, or a complementary sequence thereof. Hybridization techniques are well known in the art of molecular biology. For purposes of illustration, suitable moderately stringent conditions for testing the hybridization of a polynucleotide of this invention with other polynucleotides include prewashing in a solution of 5 X SSC, 0.5% SDS, 1.0 mM 10 EDTA (pH 8.0); hybridizing at 50°C-65°C, 5 X SSC, overnight; followed by washing twice at 65°C for 20 minutes with each of 2X, 0.5X and 0.2X SSC containing 0.1% SDS.

Moreover, it will be appreciated by those of ordinary skill in the art that, as a result of the degeneracy of the genetic code, there are many nucleotide sequences 15 that encode a polypeptide as described herein. Some of these polynucleotides bear minimal homology to the nucleotide sequence of any native gene. Nonetheless, polynucleotides that vary due to differences in codon usage are specifically contemplated by the present invention. Further, alleles of the genes comprising the polynucleotide sequences provided herein are within the scope of the present invention. 20 Alleles are endogenous genes that are altered as a result of one or more mutations, such as deletions, additions and/or substitutions of nucleotides. The resulting mRNA and protein may, but need not, have an altered structure or function. Alleles may be identified using standard techniques (such as hybridization, amplification and/or database sequence comparison).

25 PROBES AND PRIMERS

In other embodiments of the present invention, the polynucleotide sequences provided herein can be advantageously used as probes or primers for nucleic acid hybridization. As such, it is contemplated that nucleic acid segments that comprise a sequence region of at least about 15 nucleotide long contiguous sequence that has the 30 same sequence as, or is complementary to, a 15 nucleotide long contiguous sequence

disclosed herein will find particular utility. Longer contiguous identical or complementary sequences, *e.g.*, those of about 20, 30, 40, 50, 100, 200, 500, 1000 (including all intermediate lengths) and even up to full length sequences will also be of use in certain embodiments.

5 The ability of such nucleic acid probes to specifically hybridize to a sequence of interest will enable them to be of use in detecting the presence of complementary sequences in a given sample. However, other uses are also envisioned, such as the use of the sequence information for the preparation of mutant species primers, or primers for use in preparing other genetic constructions.

10 Polynucleotide molecules having sequence regions consisting of contiguous nucleotide stretches of 10-14, 15-20, 30, 50, or even of 100-200 nucleotides or so (including intermediate lengths as well), identical or complementary to a polynucleotide sequence disclosed herein, are particularly contemplated as hybridization probes for use in, *e.g.*, Southern and Northern blotting. This would allow
15 a gene product, or fragment thereof, to be analyzed, both in diverse cell types and also in various bacterial cells. The total size of fragment, as well as the size of the complementary stretch(es), will ultimately depend on the intended use or application of the particular nucleic acid segment. Smaller fragments will generally find use in hybridization embodiments, wherein the length of the contiguous complementary
20 region may be varied, such as between about 15 and about 100 nucleotides, but larger contiguous complementarity stretches may be used, according to the length complementary sequences one wishes to detect.

 The use of a hybridization probe of about 15-25 nucleotides in length allows the formation of a duplex molecule that is both stable and selective. Molecules
25 having contiguous complementary sequences over stretches greater than 15 bases in length are generally preferred, though, in order to increase stability and selectivity of the hybrid, and thereby improve the quality and degree of specific hybrid molecules obtained. One will generally prefer to design nucleic acid molecules having gene-complementary stretches of 15 to 25 contiguous nucleotides, or even longer where
30 desired.

Hybridization probes may be selected from any portion of any of the sequences disclosed herein. All that is required is to review the sequence set forth in SEQ ID NO: 1-38, 42-204, 205, 207 and 210-290, or to any continuous portion of the sequence, from about 15-25 nucleotides in length up to and including the full length 5 sequence, that one wishes to utilize as a probe or primer. The choice of probe and primer sequences may be governed by various factors. For example, one may wish to employ primers from towards the termini of the total sequence.

Small polynucleotide segments or fragments may be readily prepared by, for example, directly synthesizing the fragment by chemical means, as is commonly 10 practiced using an automated oligonucleotide synthesizer. Also, fragments may be obtained by application of nucleic acid reproduction technology, such as the PCR™ technology of U. S. Patent 4,683,202 (incorporated herein by reference), by introducing selected sequences into recombinant vectors for recombinant production, and by other recombinant DNA techniques generally known to those of skill in the art of molecular 15 biology.

The nucleotide sequences of the invention may be used for their ability to selectively form duplex molecules with complementary stretches of the entire gene or gene fragments of interest. Depending on the application envisioned, one will typically desire to employ varying conditions of hybridization to achieve varying degrees of 20 selectivity of probe towards target sequence. For applications requiring high selectivity, one will typically desire to employ relatively stringent conditions to form the hybrids, e.g., one will select relatively low salt and/or high temperature conditions, such as provided by a salt concentration of from about 0.02 M to about 0.15 M salt at temperatures of from about 50°C to about 70°C. Such selective conditions tolerate 25 little, if any, mismatch between the probe and the template or target strand, and would be particularly suitable for isolating related sequences.

Of course, for some applications, for example, where one desires to prepare mutants employing a mutant primer strand hybridized to an underlying template, less stringent (reduced stringency) hybridization conditions will typically be 30 needed in order to allow formation of the heteroduplex. In these circumstances, one may desire to employ salt conditions such as those of from about 0.15 M to about 0.9 M

salt, at temperatures ranging from about 20°C to about 55°C. Cross-hybridizing species can thereby be readily identified as positively hybridizing signals with respect to control hybridizations. In any case, it is generally appreciated that conditions can be rendered more stringent by the addition of increasing amounts of formamide, which serves to 5 destabilize the hybrid duplex in the same manner as increased temperature. Thus, hybridization conditions can be readily manipulated, and thus will generally be a method of choice depending on the desired results.

POLYNUCLEOTIDE IDENTIFICATION AND CHARACTERIZATION

Polynucleotides may be identified, prepared and/or manipulated using 10 any of a variety of well established techniques. For example, a polynucleotide may be identified, as described in more detail below, by screening a microarray of cDNAs for tumor-associated expression (*i.e.*, expression that is at least two fold greater in a tumor than in normal tissue, as determined using a representative assay provided herein). Such screens may be performed, for example, using a Synteni microarray (Palo Alto, 15 CA) according to the manufacturer's instructions (and essentially as described by Schena et al., *Proc. Natl. Acad. Sci. USA* 93:10614-10619, 1996 and Heller et al., *Proc. Natl. Acad. Sci. USA* 94:2150-2155, 1997). Alternatively, polynucleotides may be amplified from cDNA prepared from cells expressing the proteins described herein, such as breast tumor cells. Such polynucleotides may be amplified via polymerase 20 chain reaction (PCR). For this approach, sequence-specific primers may be designed based on the sequences provided herein, and may be purchased or synthesized.

An amplified portion of a polynucleotide of the present invention may be used to isolate a full length gene from a suitable library (*e.g.*, a breast tumor cDNA library) using well known techniques. Within such techniques, a library (cDNA or 25 genomic) is screened using one or more polynucleotide probes or primers suitable for amplification. Preferably, a library is size-selected to include larger molecules. Random primed libraries may also be preferred for identifying 5' and upstream regions of genes. Genomic libraries are preferred for obtaining introns and extending 5' sequences.

For hybridization techniques, a partial sequence may be labeled (e.g., by nick-translation or end-labeling with ^{32}P) using well known techniques. A bacterial or bacteriophage library is then generally screened by hybridizing filters containing denatured bacterial colonies (or lawns containing phage plaques) with the labeled probe
5 (see Sambrook et al., *Molecular Cloning: A Laboratory Manual*, Cold Spring Harbor Laboratories, Cold Spring Harbor, NY, 1989). Hybridizing colonies or plaques are selected and expanded, and the DNA is isolated for further analysis. cDNA clones may be analyzed to determine the amount of additional sequence by, for example, PCR using a primer from the partial sequence and a primer from the vector. Restriction maps and
10 partial sequences may be generated to identify one or more overlapping clones. The complete sequence may then be determined using standard techniques, which may involve generating a series of deletion clones. The resulting overlapping sequences can then assembled into a single contiguous sequence. A full length cDNA molecule can be generated by ligating suitable fragments, using well known techniques.

15 Alternatively, there are numerous amplification techniques for obtaining a full length coding sequence from a partial cDNA sequence. Within such techniques, amplification is generally performed via PCR. Any of a variety of commercially available kits may be used to perform the amplification step. Primers may be designed using, for example, software well known in the art. Primers are preferably 22-30
20 nucleotides in length, have a GC content of at least 50% and anneal to the target sequence at temperatures of about 68°C to 72°C. The amplified region may be sequenced as described above, and overlapping sequences assembled into a contiguous sequence.

One such amplification technique is inverse PCR (see Triglia et al., *Nucl.
25 Acids Res.* 16:8186, 1988), which uses restriction enzymes to generate a fragment in the known region of the gene. The fragment is then circularized by intramolecular ligation and used as a template for PCR with divergent primers derived from the known region. Within an alternative approach, sequences adjacent to a partial sequence may be retrieved by amplification with a primer to a linker sequence and a primer specific to a
30 known region. The amplified sequences are typically subjected to a second round of amplification with the same linker primer and a second primer specific to the known

region. A variation on this procedure, which employs two primers that initiate extension in opposite directions from the known sequence, is described in WO 96/38591. Another such technique is known as "rapid amplification of cDNA ends" or RACE. This technique involves the use of an internal primer and an external primer, 5 which hybridizes to a polyA region or vector sequence, to identify sequences that are 5' and 3' of a known sequence. Additional techniques include capture PCR (Lagerstrom et al., *PCR Methods Applic.* 1:111-19, 1991) and walking PCR (Parker et al., *Nucl. Acids. Res.* 19:3055-60, 1991). Other methods employing amplification may also be employed to obtain a full length cDNA sequence.

10 In certain instances, it is possible to obtain a full length cDNA sequence by analysis of sequences provided in an expressed sequence tag (EST) database, such as that available from GenBank. Searches for overlapping ESTs may generally be performed using well known programs (e.g., NCBI BLAST searches), and such ESTs may be used to generate a contiguous full length sequence. Full length DNA sequences 15 may also be obtained by analysis of genomic fragments.

POLYNUCLEOTIDE EXPRESSION IN HOST CELLS

In other embodiments of the invention, polynucleotide sequences or fragments thereof which encode polypeptides of the invention, or fusion proteins or functional equivalents thereof, may be used in recombinant DNA molecules to direct 20 expression of a polypeptide in appropriate host cells. Due to the inherent degeneracy of the genetic code, other DNA sequences that encode substantially the same or a functionally equivalent amino acid sequence may be produced and these sequences may be used to clone and express a given polypeptide.

As will be understood by those of skill in the art, it may be advantageous 25 in some instances to produce polypeptide-encoding nucleotide sequences possessing non-naturally occurring codons. For example, codons preferred by a particular prokaryotic or eukaryotic host can be selected to increase the rate of protein expression or to produce a recombinant RNA transcript having desirable properties, such as a half-life which is longer than that of a transcript generated from the naturally occurring 30 sequence.

Moreover, the polynucleotide sequences of the present invention can be engineered using methods generally known in the art in order to alter polypeptide encoding sequences for a variety of reasons, including but not limited to, alterations which modify the cloning, processing, and/or expression of the gene product. For 5 example, DNA shuffling by random fragmentation and PCR reassembly of gene fragments and synthetic oligonucleotides may be used to engineer the nucleotide sequences. In addition, site-directed mutagenesis may be used to insert new restriction sites, alter glycosylation patterns, change codon preference, produce splice variants, or introduce mutations, and so forth.

10 In another embodiment of the invention, natural, modified, or recombinant nucleic acid sequences may be ligated to a heterologous sequence to encode a fusion protein. For example, to screen peptide libraries for inhibitors of polypeptide activity, it may be useful to encode a chimeric protein that can be recognized by a commercially available antibody. A fusion protein may also be 15 engineered to contain a cleavage site located between the polypeptide-encoding sequence and the heterologous protein sequence, so that the polypeptide may be cleaved and purified away from the heterologous moiety.

Sequences encoding a desired polypeptide may be synthesized, in whole or in part, using chemical methods well known in the art (see Caruthers, M. H. et al. 20 (1980) *Nucl. Acids Res. Symp. Ser.* 215-223, Horn, T. et al. (1980) *Nucl. Acids Res. Symp. Ser.* 225-232). Alternatively, the protein itself may be produced using chemical methods to synthesize the amino acid sequence of a polypeptide, or a portion thereof. For example, peptide synthesis can be performed using various solid-phase techniques (Roberge, J. Y. et al. (1995) *Science* 269:202-204) and automated synthesis may be 25 achieved, for example, using the ABI 431A Peptide Synthesizer (Perkin Elmer, Palo Alto, CA).

A newly synthesized peptide may be substantially purified by preparative high performance liquid chromatography (e.g., Creighton, T. (1983) Proteins, Structures and Molecular Principles, WH Freeman and Co., New York, N.Y.) 30 or other comparable techniques available in the art. The composition of the synthetic peptides may be confirmed by amino acid analysis or sequencing (e.g., the Edman

degradation procedure). Additionally, the amino acid sequence of a polypeptide, or any part thereof, may be altered during direct synthesis and/or combined using chemical methods with sequences from other proteins, or any part thereof, to produce a variant polypeptide.

5 In order to express a desired polypeptide, the nucleotide sequences encoding the polypeptide, or functional equivalents, may be inserted into appropriate expression vector, i.e., a vector which contains the necessary elements for the transcription and translation of the inserted coding sequence. Methods which are well known to those skilled in the art may be used to construct expression vectors containing
10 sequences encoding a polypeptide of interest and appropriate transcriptional and translational control elements. These methods include in vitro recombinant DNA techniques, synthetic techniques, and in vivo genetic recombination. Such techniques are described in Sambrook, J. et al. (1989) Molecular Cloning, A Laboratory Manual, Cold Spring Harbor Press, Plainview, N.Y., and Ausubel, F. M. et al. (1989) Current
15 Protocols in Molecular Biology, John Wiley & Sons, New York. N.Y.

A variety of expression vector/host systems may be utilized to contain and express polynucleotide sequences. These include, but are not limited to, microorganisms such as bacteria transformed with recombinant bacteriophage, plasmid, or cosmid DNA expression vectors; yeast transformed with yeast expression vectors;
20 insect cell systems infected with virus expression vectors (e.g., baculovirus); plant cell systems transformed with virus expression vectors (e.g., cauliflower mosaic virus, CaMV; tobacco mosaic virus, TMV) or with bacterial expression vectors (e.g., Ti or pBR322 plasmids); or animal cell systems.

The "control elements" or "regulatory sequences" present in an
25 expression vector are those non-translated regions of the vector--enhancers, promoters, 5' and 3' untranslated regions--which interact with host cellular proteins to carry out transcription and translation. Such elements may vary in their strength and specificity. Depending on the vector system and host utilized, any number of suitable transcription and translation elements, including constitutive and inducible promoters, may be used.
30 For example, when cloning in bacterial systems, inducible promoters such as the hybrid lacZ promoter of the PBLUESCRIPT phagemid (Stratagene, La Jolla, Calif.) or

PSPORT1 plasmid (Gibco BRL, Gaithersburg, MD) and the like may be used. In mammalian cell systems, promoters from mammalian genes or from mammalian viruses are generally preferred. If it is necessary to generate a cell line that contains multiple copies of the sequence encoding a polypeptide, vectors based on SV40 or EBV 5 may be advantageously used with an appropriate selectable marker.

In bacterial systems, a number of expression vectors may be selected depending upon the use intended for the expressed polypeptide. For example, when large quantities are needed, for example for the induction of antibodies, vectors which direct high level expression of fusion proteins that are readily purified may be used: 10 Such vectors include, but are not limited to, the multifunctional *E. coli* cloning and expression vectors such as BLUESCRIPT (Stratagene), in which the sequence encoding the polypeptide of interest may be ligated into the vector in frame with sequences for the amino-terminal Met and the subsequent 7 residues of .beta.-galactosidase so that a hybrid protein is produced; pIN vectors (Van Heeke, G. and S. M. Schuster (1989) *J. 15 Biol. Chem.* 264:5503-5509); and the like. pGEX Vectors (Promega, Madison, Wis.) may also be used to express foreign polypeptides as fusion proteins with glutathione S-transferase (GST). In general, such fusion proteins are soluble and can easily be purified from lysed cells by adsorption to glutathione-agarose beads followed by elution in the presence of free glutathione. Proteins made in such systems may be designed to include 20 heparin, thrombin, or factor XA protease cleavage sites so that the cloned polypeptide of interest can be released from the GST moiety at will.

In the yeast, *Saccharomyces cerevisiae*, a number of vectors containing constitutive or inducible promoters such as alpha factor, alcohol oxidase, and PGH may be used. For reviews, see Ausubel et al. (supra) and Grant et al. (1987) *Methods 25 Enzymol.* 153:516-544.

In cases where plant expression vectors are used, the expression of sequences encoding polypeptides may be driven by any of a number of promoters. For example, viral promoters such as the 35S and 19S promoters of CaMV may be used alone or in combination with the omega leader sequence from TMV (Takamatsu, N. 30 (1987) *EMBO J.* 6:307-311. Alternatively, plant promoters such as the small subunit of RUBISCO or heat shock promoters may be used (Coruzzi, G. et al. (1984) *EMBO J.*

3:1671-1680; Broglie, R. et al. (1984) *Science* 224:838-843; and Winter, J. et al. (1991) *Results Probl. Cell Differ.* 17:85-105). These constructs can be introduced into plant cells by direct DNA transformation or pathogen-mediated transfection. Such techniques are described in a number of generally available reviews (see, for example, Hobbs, S. or 5 Murry, L. E. in McGraw Hill Yearbook of Science and Technology (1992) McGraw Hill, New York, N.Y.; pp. 191-196).

An insect system may also be used to express a polypeptide of interest. For example, in one such system, *Autographa californica* nuclear polyhedrosis virus (AcNPV) is used as a vector to express foreign genes in *Spodoptera frugiperda* cells or 10 in *Trichoplusia* larvae. The sequences encoding the polypeptide may be cloned into a non-essential region of the virus, such as the polyhedrin gene, and placed under control of the polyhedrin promoter. Successful insertion of the polypeptide-encoding sequence will render the polyhedrin gene inactive and produce recombinant virus lacking coat protein. The recombinant viruses may then be used to infect, for example, *S. frugiperda* 15 cells or *Trichoplusia* larvae in which the polypeptide of interest may be expressed (Engelhard, E. K. et al. (1994) *Proc. Natl. Acad. Sci.* 91 :3224-3227).

In mammalian host cells, a number of viral-based expression systems are generally available. For example, in cases where an adenovirus is used as an expression vector, sequences encoding a polypeptide of interest may be ligated into an adenovirus 20 transcription/translation complex consisting of the late promoter and tripartite leader sequence. Insertion in a non-essential E1 or E3 region of the viral genome may be used to obtain a viable virus which is capable of expressing the polypeptide in infected host cells (Logan, J. and Shenk, T. (1984) *Proc. Natl. Acad. Sci.* 81:3655-3659). In addition, transcription enhancers, such as the Rous sarcoma virus (RSV) enhancer, may be used 25 to increase expression in mammalian host cells.

Specific initiation signals may also be used to achieve more efficient translation of sequences encoding a polypeptide of interest. Such signals include the ATG initiation codon and adjacent sequences. In cases where sequences encoding the polypeptide, its initiation codon, and upstream sequences are inserted into the 30 appropriate expression vector, no additional transcriptional or translational control signals may be needed. However, in cases where only coding sequence, or a portion

thereof, is inserted, exogenous translational control signals including the ATG initiation codon should be provided. Furthermore, the initiation codon should be in the correct reading frame to ensure translation of the entire insert. Exogenous translational elements and initiation codons may be of various origins, both natural and synthetic.

5 The efficiency of expression may be enhanced by the inclusion of enhancers which are appropriate for the particular cell system which is used, such as those described in the literature (Scharf, D. et al. (1994) *Results Probl. Cell Differ.* 20:125-162).

In addition, a host cell strain may be chosen for its ability to modulate the expression of the inserted sequences or to process the expressed protein in the 10 desired fashion. Such modifications of the polypeptide include, but are not limited to, acetylation, carboxylation, glycosylation, phosphorylation, lipidation, and acylation. Post-translational processing which cleaves a "pro" form of the protein may also be used to facilitate correct insertion, folding and/or function. Different host cells such as CHO, HeLa, MDCK, HEK293, and WI38, which have specific cellular machinery and 15 characteristic mechanisms for such post-translational activities, may be chosen to ensure the correct modification and processing of the foreign protein.

For long-term, high-yield production of recombinant proteins, stable expression is generally preferred. For example, cell lines which stably express a polynucleotide of interest may be transformed using expression vectors which may 20 contain viral origins of replication and/or endogenous expression elements and a selectable marker gene on the same or on a separate vector. Following the introduction of the vector, cells may be allowed to grow for 1-2 days in an enriched media before they are switched to selective media. The purpose of the selectable marker is to confer resistance to selection, and its presence allows growth and recovery of cells which 25 successfully express the introduced sequences. Resistant clones of stably transformed cells may be proliferated using tissue culture techniques appropriate to the cell type.

Any number of selection systems may be used to recover transformed cell lines. These include, but are not limited to, the herpes simplex virus thymidine kinase (Wigler, M. et al. (1977) *Cell* 11:223-32) and adenine phosphoribosyltransferase 30 (Lowy, I. et al. (1990) *Cell* 22:817-23) genes which can be employed in tk.sup.- or aprt.sup.- cells, respectively. Also, antimetabolite, antibiotic or herbicide resistance can

be used as the basis for selection; for example, dhfr which confers resistance to methotrexate (Wigler, M. et al. (1980) *Proc. Natl. Acad. Sci.* 77:3567-70); npt, which confers resistance to the aminoglycosides, neomycin and G-418 (Colbere-Garapin, F. et al (1981) *J. Mol. Biol.* 150:1-14); and als or pat, which confer resistance to 5 chlorsulfuron and phosphinotricin acetyltransferase, respectively (Murry, *supra*). Additional selectable genes have been described, for example, trpB, which allows cells to utilize indole in place of tryptophan, or hisD, which allows cells to utilize histinol in place of histidine (Hartman, S. C. and R. C. Mulligan (1988) *Proc. Natl. Acad. Sci.* 85:8047-51). Recently, the use of visible markers has gained popularity with such 10 markers as anthocyanins, beta-glucuronidase and its substrate GUS, and luciferase and its substrate luciferin, being widely used not only to identify transformants, but also to quantify the amount of transient or stable protein expression attributable to a specific vector system (Rhodes, C. A. et al. (1995) *Methods Mol. Biol.* 55:121-131).

Although the presence/absence of marker gene expression suggests that 15 the gene of interest is also present, its presence and expression may need to be confirmed. For example, if the sequence encoding a polypeptide is inserted within a marker gene sequence, recombinant cells containing sequences can be identified by the absence of marker gene function. Alternatively, a marker gene can be placed in tandem with a polypeptide-encoding sequence under the control of a single promoter. 20 Expression of the marker gene in response to induction or selection usually indicates expression of the tandem gene as well.

Alternatively, host cells which contain and express a desired polynucleotide sequence may be identified by a variety of procedures known to those of skill in the art. These procedures include, but are not limited to, DNA-DNA or DNA- 25 RNA hybridizations and protein bioassay or immunoassay techniques which include membrane, solution, or chip based technologies for the detection and/or quantification of nucleic acid or protein.

A variety of protocols for detecting and measuring the expression of polynucleotide-encoded products, using either polyclonal or monoclonal antibodies 30 specific for the product are known in the art. Examples include enzyme-linked immunosorbent assay (ELISA), radioimmunoassay (RIA), and fluorescence activated

cell sorting (FACS). A two-site, monoclonal-based immunoassay utilizing monoclonal antibodies reactive to two non-interfering epitopes on a given polypeptide may be preferred for some applications, but a competitive binding assay may also be employed. These and other assays are described, among other places, in Hampton, R. et al. (1990; 5 Serological Methods, a Laboratory Manual, APS Press, St Paul. Minn.) and Maddox, D. E. et al. (1983; *J. Exp. Med.* 158:1211-1216).

A wide variety of labels and conjugation techniques are known by those skilled in the art and may be used in various nucleic acid and amino acid assays. Means for producing labeled hybridization or PCR probes for detecting sequences related to 10 polynucleotides include oligolabeling, nick translation, end-labeling or PCR amplification using a labeled nucleotide. Alternatively, the sequences, or any portions thereof may be cloned into a vector for the production of an mRNA probe. Such vectors are known in the art, are commercially available, and may be used to synthesize RNA probes in vitro by addition of an appropriate RNA polymerase such as T7, T3, or SP6 15 and labeled nucleotides. These procedures may be conducted using a variety of commercially available kits. Suitable reporter molecules or labels, which may be used include radionuclides, enzymes, fluorescent, chemiluminescent, or chromogenic agents as well as substrates, cofactors, inhibitors, magnetic particles, and the like.

Host cells transformed with a polynucleotide sequence of interest may be 20 cultured under conditions suitable for the expression and recovery of the protein from cell culture. The protein produced by a recombinant cell may be secreted or contained intracellularly depending on the sequence and/or the vector used. As will be understood by those of skill in the art, expression vectors containing polynucleotides of the invention may be designed to contain signal sequences which direct secretion of the 25 encoded polypeptide through a prokaryotic or eukaryotic cell membrane. Other recombinant constructions may be used to join sequences encoding a polypeptide of interest to nucleotide sequence encoding a polypeptide domain which will facilitate purification of soluble proteins. Such purification facilitating domains include, but are not limited to, metal chelating peptides such as histidine-tryptophan modules that allow 30 purification on immobilized metals, protein A domains that allow purification on immobilized immunoglobulin, and the domain utilized in the FLAGS extension/affinity

purification system (Immunex Corp., Seattle, Wash.). The inclusion of cleavable linker sequences such as those specific for Factor XA or enterokinase (Invitrogen, San Diego, Calif.) between the purification domain and the encoded polypeptide may be used to facilitate purification. One such expression vector provides for expression of a fusion 5 protein containing a polypeptide of interest and a nucleic acid encoding 6 histidine residues preceding a thioredoxin or an enterokinase cleavage site. The histidine residues facilitate purification on IMIAC (immobilized metal ion affinity chromatography) as described in Porath, J. et al. (1992, *Prot. Exp. Purif.* 3:263-281) while the enterokinase cleavage site provides a means for purifying the desired polypeptide from the fusion 10 protein. A discussion of vectors which contain fusion proteins is provided in Kroll, D. J. et al. (1993; *DNA Cell Biol.* 12:441-453).

In addition to recombinant production methods, polypeptides of the invention, and fragments thereof, may be produced by direct peptide synthesis using solid-phase techniques (Merrifield J. (1963) *J. Am. Chem. Soc.* 85:2149-2154). Protein 15 synthesis may be performed using manual techniques or by automation. Automated synthesis may be achieved, for example, using Applied Biosystems 431A Peptide Synthesizer (Perkin Elmer). Alternatively, various fragments may be chemically synthesized separately and combined using chemical methods to produce the full length molecule.

20 SITE-SPECIFIC MUTAGENESIS

Site-specific mutagenesis is a technique useful in the preparation of individual peptides, or biologically functional equivalent polypeptides, through specific mutagenesis of the underlying polynucleotides that encode them. The technique, well-known to those of skill in the art, further provides a ready ability to prepare and 25 test sequence variants, for example, incorporating one or more of the foregoing considerations, by introducing one or more nucleotide sequence changes into the DNA. Site-specific mutagenesis allows the production of mutants through the use of specific oligonucleotide sequences which encode the DNA sequence of the desired mutation, as well as a sufficient number of adjacent nucleotides, to provide a primer sequence of 30 sufficient size and sequence complexity to form a stable duplex on both sides of the

deletion junction being traversed. Mutations may be employed in a selected polynucleotide sequence to improve, alter, decrease, modify, or otherwise change the properties of the polynucleotide itself, and/or alter the properties, activity, composition, stability, or primary sequence of the encoded polypeptide.

5 In certain embodiments of the present invention, the inventors contemplate the mutagenesis of the disclosed polynucleotide sequences to alter one or more properties of the encoded polypeptide, such as the antigenicity of a polypeptide vaccine. The techniques of site-specific mutagenesis are well-known in the art, and are widely used to create variants of both polypeptides and polynucleotides. For example,
10 site-specific mutagenesis is often used to alter a specific portion of a DNA molecule. In such embodiments, a primer comprising typically about 14 to about 25 nucleotides or so in length is employed, with about 5 to about 10 residues on both sides of the junction of the sequence being altered.

As will be appreciated by those of skill in the art, site-specific
15 mutagenesis techniques have often employed a phage vector that exists in both a single stranded and double stranded form. Typical vectors useful in site-directed mutagenesis include vectors such as the M13 phage. These phage are readily commercially-available and their use is generally well-known to those skilled in the art. Double-stranded plasmids are also routinely employed in site directed mutagenesis that
20 eliminates the step of transferring the gene of interest from a plasmid to a phage.

In general, site-directed mutagenesis in accordance herewith is performed by first obtaining a single-stranded vector or melting apart of two strands of a double-stranded vector that includes within its sequence a DNA sequence that encodes the desired peptide. An oligonucleotide primer bearing the desired mutated sequence is
25 prepared, generally synthetically. This primer is then annealed with the single-stranded vector, and subjected to DNA polymerizing enzymes such as *E. coli* polymerase I Klenow fragment, in order to complete the synthesis of the mutation-bearing strand. Thus, a heteroduplex is formed wherein one strand encodes the original non-mutated sequence and the second strand bears the desired mutation. This heteroduplex vector is
30 then used to transform appropriate cells, such as *E. coli* cells, and clones are selected which include recombinant vectors bearing the mutated sequence arrangement.

The preparation of sequence variants of the selected peptide-encoding DNA segments using site-directed mutagenesis provides a means of producing potentially useful species and is not meant to be limiting as there are other ways in which sequence variants of peptides and the DNA sequences encoding them may be 5 obtained. For example, recombinant vectors encoding the desired peptide sequence may be treated with mutagenic agents, such as hydroxylamine, to obtain sequence variants. Specific details regarding these methods and protocols are found in the teachings of Maloy *et al.*, 1994; Segal, 1976; Prokop and Bajpai, 1991; Kuby, 1994; and Maniatis *et al.*, 1982, each incorporated herein by reference, for that purpose.

10 As used herein, the term "oligonucleotide directed mutagenesis procedure" refers to template-dependent processes and vector-mediated propagation which result in an increase in the concentration of a specific nucleic acid molecule relative to its initial concentration, or in an increase in the concentration of a detectable signal, such as amplification. As used herein, the term "oligonucleotide directed 15 mutagenesis procedure" is intended to refer to a process that involves the template-dependent extension of a primer molecule. The term template dependent process refers to nucleic acid synthesis of an RNA or a DNA molecule wherein the sequence of the newly synthesized strand of nucleic acid is dictated by the well-known rules of complementary base pairing (see, for example, Watson, 1987). Typically, 20 vector mediated methodologies involve the introduction of the nucleic acid fragment into a DNA or RNA vector, the clonal amplification of the vector, and the recovery of the amplified nucleic acid fragment. Examples of such methodologies are provided by U. S. Patent No. 4,237,224, specifically incorporated herein by reference in its entirety.

POLYNUCLEOTIDE AMPLIFICATION TECHNIQUES

25 A number of template dependent processes are available to amplify the target sequences of interest present in a sample. One of the best known amplification methods is the polymerase chain reaction (PCRTM) which is described in detail in U.S. Patent Nos. 4,683,195, 4,683,202 and 4,800,159, each of which is incorporated herein by reference in its entirety. Briefly, in PCRTM, two primer sequences are prepared 30 which are complementary to regions on opposite complementary strands of the target

sequence. An excess of deoxynucleoside triphosphates is added to a reaction mixture along with a DNA polymerase (e.g., *Taq* polymerase). If the target sequence is present in a sample, the primers will bind to the target and the polymerase will cause the primers to be extended along the target sequence by adding on nucleotides. By raising 5 and lowering the temperature of the reaction mixture, the extended primers will dissociate from the target to form reaction products, excess primers will bind to the target and to the reaction product and the process is repeated. Preferably reverse transcription and PCR™ amplification procedure may be performed in order to quantify the amount of mRNA amplified. Polymerase chain reaction methodologies are well 10 known in the art.

Another method for amplification is the ligase chain reaction (referred to as LCR), disclosed in Eur. Pat. Appl. Publ. No. 320,308 (specifically incorporated herein by reference in its entirety). In LCR, two complementary probe pairs are prepared, and in the presence of the target sequence, each pair will bind to opposite 15 complementary strands of the target such that they abut. In the presence of a ligase, the two probe pairs will link to form a single unit. By temperature cycling, as in PCR™, bound ligated units dissociate from the target and then serve as "target sequences" for ligation of excess probe pairs. U.S. Patent No. 4,883,750, incorporated herein by reference in its entirety, describes an alternative method of amplification similar to LCR 20 for binding probe pairs to a target sequence.

Qbeta Replicase, described in PCT Int'l. Pat. Appl. Publ. No. PCT/US87/00880, incorporated herein by reference in its entirety, may also be used as still another amplification method in the present invention. In this method, a replicative sequence of RNA that has a region complementary to that of a target is added to a 25 sample in the presence of an RNA polymerase. The polymerase will copy the replicative sequence that can then be detected.

An isothermal amplification method, in which restriction endonucleases and ligases are used to achieve the amplification of target molecules that contain nucleotide 5'-[α -thio]triphosphates in one strand of a restriction site (Walker *et al.*, 30 1992, incorporated herein by reference in its entirety), may also be useful in the amplification of nucleic acids in the present invention.

Strand Displacement Amplification (SDA) is another method of carrying out isothermal amplification of nucleic acids which involves multiple rounds of strand displacement and synthesis, *i.e.* nick translation. A similar method, called Repair Chain Reaction (RCR) is another method of amplification which may be useful in the present invention and is involves annealing several probes throughout a region targeted for amplification, followed by a repair reaction in which only two of the four bases are present. The other two bases can be added as biotinylated derivatives for easy detection. A similar approach is used in SDA.

Sequences can also be detected using a cyclic probe reaction (CPR). In CPR, a probe having a 3' and 5' sequences of non-target DNA and an internal or "middle" sequence of the target protein specific RNA is hybridized to DNA which is present in a sample. Upon hybridization, the reaction is treated with RNaseH, and the products of the probe are identified as distinctive products by generating a signal that is released after digestion. The original template is annealed to another cycling probe and the reaction is repeated. Thus, CPR involves amplifying a signal generated by hybridization of a probe to a target gene specific expressed nucleic acid.

Still other amplification methods described in Great Britain Pat. Appl. No. 2 202 328, and in PCT Intl. Pat. Appl. Publ. No. PCT/US89/01025, each of which is incorporated herein by reference in its entirety, may be used in accordance with the present invention. In the former application, "modified" primers are used in a PCR-like, template and enzyme dependent synthesis. The primers may be modified by labeling with a capture moiety (*e.g.*, biotin) and/or a detector moiety (*e.g.*, enzyme). In the latter application, an excess of labeled probes is added to a sample. In the presence of the target sequence, the probe binds and is cleaved catalytically. After cleavage, the target sequence is released intact to be bound by excess probe. Cleavage of the labeled probe signals the presence of the target sequence.

Other nucleic acid amplification procedures include transcription-based amplification systems (TAS) (Kwoh *et al.*, 1989; PCT Intl. Pat. Appl. Publ. No. WO 88/10315, incorporated herein by reference in its entirety), including nucleic acid sequence based amplification (NASBA) and 3SR. In NASBA, the nucleic acids can be prepared for amplification by standard phenol/chloroform extraction, heat denaturation

of a sample, treatment with lysis buffer and minispin columns for isolation of DNA and RNA or guanidinium chloride extraction of RNA. These amplification techniques involve annealing a primer that has sequences specific to the target sequence. Following polymerization, DNA/RNA hybrids are digested with RNase H while double stranded DNA molecules are heat-denatured again. In either case the single stranded DNA is made fully double stranded by addition of second target-specific primer, followed by polymerization. The double stranded DNA molecules are then multiply transcribed by a polymerase such as T7 or SP6. In an isothermal cyclic reaction, the RNAs are reverse transcribed into DNA, and transcribed once again with a polymerase such as T7 or SP6. The resulting products, whether truncated or complete, indicate target-specific sequences.

Eur. Pat. Appl. Publ. No. 329,822, incorporated herein by reference in its entirety, disclose a nucleic acid amplification process involving cyclically synthesizing single-stranded RNA ("ssRNA"), ssDNA, and double-stranded DNA (dsDNA), which may be used in accordance with the present invention. The ssRNA is a first template for a first primer oligonucleotide, which is elongated by reverse transcriptase (RNA-dependent DNA polymerase). The RNA is then removed from resulting DNA:RNA duplex by the action of ribonuclease H (RNase H, an RNase specific for RNA in a duplex with either DNA or RNA). The resultant ssDNA is a second template for a second primer, which also includes the sequences of an RNA polymerase promoter (exemplified by T7 RNA polymerase) 5' to its homology to its template. This primer is then extended by DNA polymerase (exemplified by the large "Klenow" fragment of *E. coli* DNA polymerase I), resulting as a double-stranded DNA ("dsDNA") molecule, having a sequence identical to that of the original RNA between the primers and having additionally, at one end, a promoter sequence. This promoter sequence can be used by the appropriate RNA polymerase to make many RNA copies of the DNA. These copies can then re-enter the cycle leading to very swift amplification. With proper choice of enzymes, this amplification can be done isothermally without addition of enzymes at each cycle. Because of the cyclical nature of this process, the starting sequence can be chosen to be in the form of either DNA or RNA.

PCT Intl. Pat. Appl. Publ. No. WO 89/06700, incorporated herein by reference in its entirety, disclose a nucleic acid sequence amplification scheme based on the hybridization of a promoter/primer sequence to a target single-stranded DNA ("ssDNA") followed by transcription of many RNA copies of the sequence. This 5 scheme is not cyclic; i.e. new templates are not produced from the resultant RNA transcripts. Other amplification methods include "RACE" (Frohman, 1990), and "one-sided PCR" (Ohara, 1989) which are well-known to those of skill in the art.

Methods based on ligation of two (or more) oligonucleotides in the presence of nucleic acid having the sequence of the resulting "di-oligonucleotide", 10 thereby amplifying the di-oligonucleotide (Wu and Dean, 1996, incorporated herein by reference in its entirety), may also be used in the amplification of DNA sequences of the present invention.

BIOLOGICAL FUNCTIONAL EQUIVALENTS

Modification and changes may be made in the structure of the 15 polynucleotides and polypeptides of the present invention and still obtain a functional molecule that encodes a polypeptide with desirable characteristics. As mentioned above, it is often desirable to introduce one or more mutations into a specific polynucleotide sequence. In certain circumstances, the resulting encoded polypeptide sequence is altered by this mutation, or in other cases, the sequence of the polypeptide 20 is unchanged by one or more mutations in the encoding polynucleotide.

When it is desirable to alter the amino acid sequence of a polypeptide to create an equivalent, or even an improved, second-generation molecule, the amino acid changes may be achieved by changing one or more of the codons of the encoding DNA sequence, according to Table 1.

25 For example, certain amino acids may be substituted for other amino acids in a protein structure without appreciable loss of interactive binding capacity with structures such as, for example, antigen-binding regions of antibodies or binding sites on substrate molecules. Since it is the interactive capacity and nature of a protein that defines that protein's biological functional activity, certain amino acid sequence 30 substitutions can be made in a protein sequence, and, of course, its underlying DNA

coding sequence, and nevertheless obtain a protein with like properties. It is thus contemplated by the inventors that various changes may be made in the peptide sequences of the disclosed compositions, or corresponding DNA sequences which encode said peptides without appreciable loss of their biological utility or activity.

5

TABLE 1

Amino Acids		Codons						
Alanine	Ala	A	GCA	GCC	GCG	GCU		
Cysteine	Cys	C	UGC	UGU				
Aspartic acid	Asp	D	GAC	GAU				
Glutamic acid	Glu	E	GAA	GAG				
Phenylalanine	Phe	F	UUC	UUU				
Glycine	Gly	G	GGA	GGC	GGG	GGU		
Histidine	His	H	CAC	CAU				
Isoleucine	Ile	I	AUA	AUC	AUU			
Lysine	Lys	K	AAA	AAG				
Leucine	Leu	L	UUA	UUG	CUA	CUC	CUG	CUU
Methionine	Met	M	AUG					
Asparagine	Asn	N	AAC	AAU				
Proline	Pro	P	CCA	CCC	CCG	CCU		
Glutamine	Gln	Q	CAA	CAG				
Arginine	Arg	R	AGA	AGG	CGA	CGC	CGG	CGU
Serine	Ser	S	AGC	AGU	UCA	UCC	UCG	UCU
Threonine	Thr	T	ACA	ACC	ACG	ACU		
Valine	Val	V	GUA	GUC	GUG	GUU		
Tryptophan	Trp	W	UGG					
Tyrosine	Tyr	Y	UAC	UAU				

In making such changes, the hydropathic index of amino acids may be considered. The importance of the hydropathic amino acid index in conferring interactive biologic function on a protein is generally understood in the art (Kyte and Doolittle, 1982, incorporated herein by reference). It is accepted that the relative

Doolittle, 1982, incorporated herein by reference). It is accepted that the relative hydrophobic character of the amino acid contributes to the secondary structure of the resultant protein, which in turn defines the interaction of the protein with other molecules, for example, enzymes, substrates, receptors, DNA, antibodies, antigens, and
5 the like. Each amino acid has been assigned a hydrophobic index on the basis of its hydrophobicity and charge characteristics (Kyte and Doolittle, 1982). These values are: isoleucine (+4.5); valine (+4.2); leucine (+3.8); phenylalanine (+2.8); cysteine/cystine (+2.5); methionine (+1.9); alanine (+1.8); glycine (-0.4); threonine (-0.7); serine (-0.8); tryptophan (-0.9); tyrosine (-1.3); proline (-1.6); histidine (-3.2); glutamate (-3.5);
10 glutamine (-3.5); aspartate (-3.5); asparagine (-3.5); lysine (-3.9); and arginine (-4.5).

It is known in the art that certain amino acids may be substituted by other amino acids having a similar hydrophobic index or score and still result in a protein with similar biological activity, *i.e.* still obtain a biological functionally equivalent protein. In making such changes, the substitution of amino acids whose
15 hydrophobic indices are within ± 2 is preferred, those within ± 1 are particularly preferred, and those within ± 0.5 are even more particularly preferred. It is also understood in the art that the substitution of like amino acids can be made effectively on the basis of hydrophilicity. U. S. Patent 4,554,101 (specifically incorporated herein by reference in its entirety), states that the greatest local average hydrophilicity of a
20 protein, as governed by the hydrophilicity of its adjacent amino acids, correlates with a biological property of the protein.

As detailed in U. S. Patent 4,554,101, the following hydrophilicity values have been assigned to amino acid residues: arginine (+3.0); lysine (+3.0); aspartate ($+3.0 \pm 1$); glutamate ($+3.0 \pm 1$); serine (+0.3); asparagine (+0.2); glutamine
25 (+0.2); glycine (0); threonine (-0.4); proline (-0.5 ± 1); alanine (-0.5); histidine (-0.5); cysteine (-1.0); methionine (-1.3); valine (-1.5); leucine (-1.8); isoleucine (-1.8); tyrosine (-2.3); phenylalanine (-2.5); tryptophan (-3.4). It is understood that an amino acid can be substituted for another having a similar hydrophilicity value and still obtain a biologically equivalent, and in particular, an immunologically equivalent protein. In
30 such changes, the substitution of amino acids whose hydrophilicity values are within ± 2

is preferred, those within ± 1 are particularly preferred, and those within ± 0.5 are even more particularly preferred.

As outlined above, amino acid substitutions are generally therefore based on the relative similarity of the amino acid side-chain substituents, for example, their 5 hydrophobicity, hydrophilicity, charge, size, and the like. Exemplary substitutions that take various of the foregoing characteristics into consideration are well known to those of skill in the art and include: arginine and lysine; glutamate and aspartate; serine and threonine; glutamine and asparagine; and valine, leucine and isoleucine.

In addition, any polynucleotide may be further modified to increase 10 stability *in vivo*. Possible modifications include, but are not limited to, the addition of flanking sequences at the 5' and/or 3' ends; the use of phosphorothioate or 2' O-methyl rather than phosphodiesterase linkages in the backbone; and/or the inclusion of nontraditional bases such as inosine, queosine and wybutosine, as well as acetyl-methyl-, thio- and other modified forms of adenine, cytidine, guanine, thymine and 15 uridine.

IN VIVO POLYNUCLEOTIDE DELIVERY TECHNIQUES

In additional embodiments, genetic constructs comprising one or more of the polynucleotides of the invention are introduced into cells *in vivo*. This may be achieved using any of a variety of well known approaches, several of which are outlined 20 below for the purpose of illustration.

1. ADENOVIRUS

One of the preferred methods for *in vivo* delivery of one or more nucleic acid sequences involves the use of an adenovirus expression vector. "Adenovirus expression vector" is meant to include those constructs containing adenovirus sequences 25 sufficient to (a) support packaging of the construct and (b) to express a polynucleotide that has been cloned therein in a sense or antisense orientation. Of course, in the context of an antisense construct, expression does not require that the gene product be synthesized.

The expression vector comprises a genetically engineered form of an adenovirus. Knowledge of the genetic organization of adenovirus, a 36 kb, linear, double-stranded DNA virus, allows substitution of large pieces of adenoviral DNA with foreign sequences up to 7 kb (Grunhaus and Horwitz, 1992). In contrast to retrovirus, 5 the adenoviral infection of host cells does not result in chromosomal integration because adenoviral DNA can replicate in an episomal manner without potential genotoxicity. Also, adenoviruses are structurally stable, and no genome rearrangement has been detected after extensive amplification. Adenovirus can infect virtually all epithelial cells regardless of their cell cycle stage. So far, adenoviral infection appears to be 10 linked only to mild disease such as acute respiratory disease in humans.

Adenovirus is particularly suitable for use as a gene transfer vector because of its mid-sized genome, ease of manipulation, high titer, wide target-cell range and high infectivity. Both ends of the viral genome contain 100-200 base pair inverted repeats (ITRs), which are *cis* elements necessary for viral DNA replication and 15 packaging. The early (E) and late (L) regions of the genome contain different transcription units that are divided by the onset of viral DNA replication. The E1 region (E1A and E1B) encodes proteins responsible for the regulation of transcription of the viral genome and a few cellular genes. The expression of the E2 region (E2A and E2B) results in the synthesis of the proteins for viral DNA replication. These proteins are 20 involved in DNA replication, late gene expression and host cell shut-off (Renan, 1990). The products of the late genes, including the majority of the viral capsid proteins, are expressed only after significant processing of a single primary transcript issued by the major late promoter (MLP). The MLP, (located at 16.8 m.u.) is particularly efficient during the late phase of infection, and all the mRNA's issued from this promoter possess 25 a 5'-tripartite leader (TPL) sequence which makes them preferred mRNA's for translation.

In a current system, recombinant adenovirus is generated from homologous recombination between shuttle vector and provirus vector. Due to the possible recombination between two proviral vectors, wild-type adenovirus may be 30 generated from this process. Therefore, it is critical to isolate a single clone of virus from an individual plaque and examine its genomic structure.

Generation and propagation of the current adenovirus vectors, which are replication deficient, depend on a unique helper cell line, designated 293, which was transformed from human embryonic kidney cells by Ad5 DNA fragments and constitutively expresses E1 proteins (Graham *et al.*, 1977). Since the E3 region is dispensable from the adenovirus genome (Jones and Shenk, 1978), the current adenovirus vectors, with the help of 293 cells, carry foreign DNA in either the E1, the D3 or both regions (Graham and Prevec, 1991). In nature, adenovirus can package approximately 105% of the wild-type genome (Ghosh-Choudhury *et al.*, 1987), providing capacity for about 2 extra kB of DNA. Combined with the approximately 5.5 kB of DNA that is replaceable in the E1 and E3 regions, the maximum capacity of the current adenovirus vector is under 7.5 kB, or about 15% of the total length of the vector. More than 80% of the adenovirus viral genome remains in the vector backbone and is the source of vector-borne cytotoxicity. Also, the replication deficiency of the E1-deleted virus is incomplete. For example, leakage of viral gene expression has been observed with the currently available vectors at high multiplicities of infection (MOI) (Mulligan, 1993).

Helper cell lines may be derived from human cells such as human embryonic kidney cells, muscle cells, hematopoietic cells or other human embryonic mesenchymal or epithelial cells. Alternatively, the helper cells may be derived from the cells of other mammalian species that are permissive for human adenovirus. Such cells include, *e.g.*, Vero cells or other monkey embryonic mesenchymal or epithelial cells. As stated above, the currently preferred helper cell line is 293.

Recently, Racher *et al.* (1995) disclosed improved methods for culturing 293 cells and propagating adenovirus. In one format, natural cell aggregates are grown by inoculating individual cells into 1 liter siliconized spinner flasks (Techne, Cambridge, UK) containing 100-200 ml of medium. Following stirring at 40 rpm, the cell viability is estimated with trypan blue. In another format, Fibra-Cel microcarriers (Bibby Sterlin, Stone, UK) (5 g/l) is employed as follows. A cell inoculum, resuspended in 5 ml of medium, is added to the carrier (50 ml) in a 250 ml Erlenmeyer flask and left stationary, with occasional agitation, for 1 to 4 h. The medium is then replaced with 50 ml of fresh medium and shaking initiated. For virus production, cells

are allowed to grow to about 80% confluence, after which time the medium is replaced (to 25% of the final volume) and adenovirus added at an MOI of 0.05. Cultures are left stationary overnight, following which the volume is increased to 100% and shaking commenced for another 72 h.

5 Other than the requirement that the adenovirus vector be replication defective, or at least conditionally defective, the nature of the adenovirus vector is not believed to be crucial to the successful practice of the invention. The adenovirus may be of any of the 42 different known serotypes or subgroups A-F. Adenovirus type 5 of subgroup C is the preferred starting material in order to obtain a conditional replication-
10 defective adenovirus vector for use in the present invention, since Adenovirus type 5 is a human adenovirus about which a great deal of biochemical and genetic information is known, and it has historically been used for most constructions employing adenovirus as a vector.

As stated above, the typical vector according to the present invention is
15 replication defective and will not have an adenovirus E1 region. Thus, it will be most convenient to introduce the polynucleotide encoding the gene of interest at the position from which the E1-coding sequences have been removed. However, the position of insertion of the construct within the adenovirus sequences is not critical to the invention. The polynucleotide encoding the gene of interest may also be inserted in lieu
20 of the deleted E3 region in E3 replacement vectors as described by Karlsson *et al.* (1986) or in the E4 region where a helper cell line or helper virus complements the E4 defect.

Adenovirus is easy to grow and manipulate and exhibits broad host range
in vitro and *in vivo*. This group of viruses can be obtained in high titers, e.g., 10⁹-10¹¹
25 plaque-forming units per ml, and they are highly infective. The life cycle of adenovirus does not require integration into the host cell genome. The foreign genes delivered by adenovirus vectors are episomal and, therefore, have low genotoxicity to host cells. No side effects have been reported in studies of vaccination with wild-type adenovirus (Couch *et al.*, 1963; Top *et al.*, 1971), demonstrating their safety and therapeutic
30 potential as *in vivo* gene transfer vectors.

Adenovirus vectors have been used in eukaryotic gene expression (Levrero *et al.*, 1991; Gomez-Foix *et al.*, 1992) and vaccine development (Grunhaus and Horwitz, 1992; Graham and Prevec, 1992). Recently, animal studies suggested that recombinant adenovirus could be used for gene therapy (Stratford-Perricaudet and 5 Perricaudet, 1991; Stratford-Perricaudet *et al.*, 1990; Rich *et al.*, 1993). Studies in administering recombinant adenovirus to different tissues include trachea instillation (Rosenfeld *et al.*, 1991; Rosenfeld *et al.*, 1992), muscle injection (Ragot *et al.*, 1993), peripheral intravenous injections (Herz and Gerard, 1993) and stereotactic inoculation into the brain (Le Gal La Salle *et al.*, 1993).

10 2. RETROVIRUSES

The retroviruses are a group of single-stranded RNA viruses characterized by an ability to convert their RNA to double-stranded DNA in infected cells by a process of reverse-transcription (Coffin, 1990). The resulting DNA then stably integrates into cellular chromosomes as a provirus and directs synthesis of viral 15 proteins. The integration results in the retention of the viral gene sequences in the recipient cell and its descendants. The retroviral genome contains three genes, gag, pol, and env that code for capsid proteins, polymerase enzyme, and envelope components, respectively. A sequence found upstream from the gag gene contains a signal for packaging of the genome into virions. Two long terminal repeat (LTR) sequences are 20 present at the 5' and 3' ends of the viral genome. These contain strong promoter and enhancer sequences and are also required for integration in the host cell genome (Coffin, 1990).

In order to construct a retroviral vector, a nucleic acid encoding one or more oligonucleotide or polynucleotide sequences of interest is inserted into the viral 25 genome in the place of certain viral sequences to produce a virus that is replication-defective. In order to produce virions, a packaging cell line containing the gag, pol, and env genes but without the LTR and packaging components is constructed (Mann *et al.*, 1983). When a recombinant plasmid containing a cDNA, together with the retroviral LTR and packaging sequences is introduced into this cell line (by calcium phosphate 30 precipitation for example), the packaging sequence allows the RNA transcript of the

recombinant plasmid to be packaged into viral particles, which are then secreted into the culture media (Nicolas and Rubenstein, 1988; Temin, 1986; Mann *et al.*, 1983). The media containing the recombinant retroviruses is then collected, optionally concentrated, and used for gene transfer. Retroviral vectors are able to infect a broad variety of cell types. However, integration and stable expression require the division of host cells (Paskind *et al.*, 1975).

A novel approach designed to allow specific targeting of retrovirus vectors was recently developed based on the chemical modification of a retrovirus by the chemical addition of lactose residues to the viral envelope. This modification could permit the specific infection of hepatocytes *via* sialoglycoprotein receptors.

A different approach to targeting of recombinant retroviruses was designed in which biotinylated antibodies against a retroviral envelope protein and against a specific cell receptor were used. The antibodies were coupled *via* the biotin components by using streptavidin (Roux *et al.*, 1989). Using antibodies against major histocompatibility complex class I and class II antigens, they demonstrated the infection of a variety of human cells that bore those surface antigens with an ecotropic virus *in vitro* (Roux *et al.*, 1989).

3. ADENO-ASSOCIATED VIRUSES

AAV (Ridgeway, 1988; Hermonat and Muzychka, 1984) is a parovirus, discovered as a contamination of adenoviral stocks. It is a ubiquitous virus (antibodies are present in 85% of the US human population) that has not been linked to any disease. It is also classified as a dependovirus, because its replication is dependent on the presence of a helper virus, such as adenovirus. Five serotypes have been isolated, of which AAV-2 is the best characterized. AAV has a single-stranded linear DNA that is encapsidated into capsid proteins VP1, VP2 and VP3 to form an icosahedral virion of 20 to 24 nm in diameter (Muzychka and McLaughlin, 1988).

The AAV DNA is approximately 4.7 kilobases long. It contains two open reading frames and is flanked by two ITRs. There are two major genes in the AAV genome: *rep* and *cap*. The *rep* gene codes for proteins responsible for viral replication, whereas *cap* codes for capsid protein VP1-3. Each ITR forms a T-shaped

hairpin structure. These terminal repeats are the only essential *cis* components of the AAV for chromosomal integration. Therefore, the AAV can be used as a vector with all viral coding sequences removed and replaced by the cassette of genes for delivery. Three viral promoters have been identified and named p5, p19, and p40, according to 5 their map position. Transcription from p5 and p19 results in production of rep proteins, and transcription from p40 produces the capsid proteins (Hermonat and Muzyczka, 1984).

There are several factors that prompted researchers to study the possibility of using rAAV as an expression vector. One is that the requirements for 10 delivering a gene to integrate into the host chromosome are surprisingly few. It is necessary to have the 145-bp ITRs, which are only 6% of the AAV genome. This leaves room in the vector to assemble a 4.5-kb DNA insertion. While this carrying capacity may prevent the AAV from delivering large genes, it is amply suited for delivering the antisense constructs of the present invention.

15 AAV is also a good choice of delivery vehicles due to its safety. There is a relatively complicated rescue mechanism: not only wild type adenovirus but also AAV genes are required to mobilize rAAV. Likewise, AAV is not pathogenic and not associated with any disease. The removal of viral coding sequences minimizes immune reactions to viral gene expression, and therefore, rAAV does not evoke an inflammatory 20 response.

4. OTHER VIRAL VECTORS AS EXPRESSION CONSTRUCTS

Other viral vectors may be employed as expression constructs in the present invention for the delivery of oligonucleotide or polynucleotide sequences to a host cell. Vectors derived from viruses such as vaccinia virus (Ridgeway, 1988; Coupar 25 *et al.*, 1988), lentiviruses, polio viruses and herpes viruses may be employed. They offer several attractive features for various mammalian cells (Friedmann, 1989; Ridgeway, 1988; Coupar *et al.*, 1988; Horwitz *et al.*, 1990).

With the recent recognition of defective hepatitis B viruses, new insight was gained into the structure-function relationship of different viral sequences. *In vitro* 30 studies showed that the virus could retain the ability for helper-dependent packaging

and reverse transcription despite the deletion of up to 80% of its genome (Horwitz *et al.*, 1990). This suggested that large portions of the genome could be replaced with foreign genetic material. The hepatotropism and persistence (integration) were particularly attractive properties for liver-directed gene transfer. Chang *et al.* (1991) 5 introduced the chloramphenicol acetyltransferase (CAT) gene into duck hepatitis B virus genome in the place of the polymerase, surface, and pre-surface coding sequences. It was cotransfected with wild-type virus into an avian hepatoma cell line. Culture media containing high titers of the recombinant virus were used to infect primary duckling hepatocytes. Stable CAT gene expression was detected for at least 24 days 10 after transfection (Chang *et al.*, 1991).

5. NON-VIRAL VECTORS

In order to effect expression of the oligonucleotide or polynucleotide sequences of the present invention, the expression construct must be delivered into a cell. This delivery may be accomplished *in vitro*, as in laboratory procedures for 15 transforming cells lines, or *in vivo* or *ex vivo*, as in the treatment of certain disease states. As described above, one preferred mechanism for delivery is *via* viral infection where the expression construct is encapsulated in an infectious viral particle.

Once the expression construct has been delivered into the cell the nucleic acid encoding the desired oligonucleotide or polynucleotide sequences may be 20 positioned and expressed at different sites. In certain embodiments, the nucleic acid encoding the construct may be stably integrated into the genome of the cell. This integration may be in the specific location and orientation *via* homologous recombination (gene replacement) or it may be integrated in a random, non-specific location (gene augmentation). In yet further embodiments, the nucleic acid may be 25 stably maintained in the cell as a separate, episomal segment of DNA. Such nucleic acid segments or "episomes" encode sequences sufficient to permit maintenance and replication independent of or in synchronization with the host cell cycle. How the expression construct is delivered to a cell and where in the cell the nucleic acid remains is dependent on the type of expression construct employed.

In certain embodiments of the invention, the expression construct comprising one or more oligonucleotide or polynucleotide sequences may simply consist of naked recombinant DNA or plasmids. Transfer of the construct may be performed by any of the methods mentioned above which physically or chemically 5 permeabilize the cell membrane. This is particularly applicable for transfer *in vitro* but it may be applied to *in vivo* use as well. Dubensky *et al.* (1984) successfully injected polyomavirus DNA in the form of calcium phosphate precipitates into liver and spleen of adult and newborn mice demonstrating active viral replication and acute infection. Benvenisty and Reshef (1986) also demonstrated that direct intraperitoneal injection of 10 calcium phosphate-precipitated plasmids results in expression of the transfected genes. It is envisioned that DNA encoding a gene of interest may also be transferred in a similar manner *in vivo* and express the gene product.

Another embodiment of the invention for transferring a naked DNA expression construct into cells may involve particle bombardment. This method 15 depends on the ability to accelerate DNA-coated microprojectiles to a high velocity allowing them to pierce cell membranes and enter cells without killing them (Klein *et al.*, 1987). Several devices for accelerating small particles have been developed. One such device relies on a high voltage discharge to generate an electrical current, which in turn provides the motive force (Yang *et al.*, 1990). The microprojectiles used have 20 consisted of biologically inert substances such as tungsten or gold beads.

Selected organs including the liver, skin, and muscle tissue of rats and mice have been bombarded *in vivo* (Yang *et al.*, 1990; Zelenin *et al.*, 1991). This may require surgical exposure of the tissue or cells, to eliminate any intervening tissue between the gun and the target organ, *i.e. ex vivo* treatment. Again, DNA encoding a 25 particular gene may be delivered *via* this method and still be incorporated by the present invention.

ANTISENSE OLIGONUCLEOTIDES

The end result of the flow of genetic information is the synthesis of protein. DNA is transcribed by polymerases into messenger RNA and translated on the 30 ribosome to yield a folded, functional protein. Thus there are several steps along the

route where protein synthesis can be inhibited. The native DNA segment coding for a polypeptide described herein, as all such mammalian DNA strands, has two strands: a sense strand and an antisense strand held together by hydrogen bonding. The messenger RNA coding for polypeptide has the same nucleotide sequence as the sense 5 DNA strand except that the DNA thymidine is replaced by uridine. Thus, synthetic antisense nucleotide sequences will bind to a mRNA and inhibit expression of the protein encoded by that mRNA.

The targeting of antisense oligonucleotides to mRNA is thus one mechanism to shut down protein synthesis, and, consequently, represents a powerful 10 and targeted therapeutic approach. For example, the synthesis of polygalacturonase and the muscarine type 2 acetylcholine receptor are inhibited by antisense oligonucleotides directed to their respective mRNA sequences (U. S. Patent 5,739,119 and U. S. Patent 5,759,829, each specifically incorporated herein by reference in its entirety). Further, examples of antisense inhibition have been demonstrated with the 15 nuclear protein cyclin, the multiple drug resistance gene (MDG1), ICAM-1, E-selectin, STK-1, striatal GABA_A receptor and human EGF (Jaskulski *et al.*, 1988; Vasanthakumar and Ahmed, 1989; Peris *et al.*, 1998; U. S. Patent 5,801,154; U. S. Patent 5,789,573; U. S. Patent 5,718,709 and U. S. Patent 5,610,288, each specifically incorporated herein by reference in its entirety). Antisense constructs have also been 20 described that inhibit and can be used to treat a variety of abnormal cellular proliferations, e.g. cancer (U. S. Patent 5,747,470; U. S. Patent 5,591,317 and U. S. Patent 5,783,683, each specifically incorporated herein by reference in its entirety).

Therefore, in exemplary embodiments, the invention provides 25 oligonucleotide sequences that comprise all, or a portion of, any sequence that is capable of specifically binding to polynucleotide sequence described herein, or a complement thereof. In one embodiment, the antisense oligonucleotides comprise DNA or derivatives thereof. In another embodiment, the oligonucleotides comprise RNA or derivatives thereof. In a third embodiment, the oligonucleotides are modified DNAs comprising a phosphorothioated modified backbone. In a fourth embodiment, the 30 oligonucleotide sequences comprise peptide nucleic acids or derivatives thereof. In each case, preferred compositions comprise a sequence region that is complementary,

and more preferably substantially-complementary, and even more preferably, completely complementary to one or more portions of polynucleotides disclosed herein.

Selection of antisense compositions specific for a given gene sequence is based upon analysis of the chosen target sequence (*i.e.* in these illustrative examples the 5 rat and human sequences) and determination of secondary structure, T_m , binding energy, relative stability, and antisense compositions were selected based upon their relative inability to form dimers, hairpins, or other secondary structures that would reduce or prohibit specific binding to the target mRNA in a host cell.

Highly preferred target regions of the mRNA, are those which are at or 10 near the AUG translation initiation codon, and those sequences which were substantially complementary to 5' regions of the mRNA. These secondary structure analyses and target site selection considerations were performed using v.4 of the OLIGO primer analysis software (Rychlik, 1997) and the BLASTN 2.0.5 algorithm software (Altschul *et al.*, 1997).

15 The use of an antisense delivery method employing a short peptide vector, termed MPG (27 residues), is also contemplated. The MPG peptide contains a hydrophobic domain derived from the fusion sequence of HIV gp41 and a hydrophilic domain from the nuclear localization sequence of SV40 T-antigen (Morris *et al.*, 1997). It has been demonstrated that several molecules of the MPG peptide coat the antisense 20 oligonucleotides and can be delivered into cultured mammalian cells in less than 1 hour with relatively high efficiency (90%). Further, the interaction with MPG strongly increases both the stability of the oligonucleotide to nuclease and the ability to cross the plasma membrane (Morris *et al.*, 1997).

RIBOZYMES

25 Although proteins traditionally have been used for catalysis of nucleic acids, another class of macromolecules has emerged as useful in this endeavor. Ribozymes are RNA-protein complexes that cleave nucleic acids in a site-specific fashion. Ribozymes have specific catalytic domains that possess endonuclease activity (Kim and Cech, 1987; Gerlach *et al.*, 1987; Forster and Symons, 1987). For example, a 30 large number of ribozymes accelerate phosphoester transfer reactions with a high degree

of specificity, often cleaving only one of several phosphoesters in an oligonucleotide substrate (Cech *et al.*, 1981; Michel and Westhof, 1990; Reinhold-Hurek and Shub, 1992). This specificity has been attributed to the requirement that the substrate bind via specific base-pairing interactions to the internal guide sequence ("IGS") of the ribozyme
5 prior to chemical reaction.

Ribozyme catalysis has primarily been observed as part of sequence-specific cleavage/ligation reactions involving nucleic acids (Joyce, 1989; Cech *et al.*, 1981). For example, U. S. Patent No. 5,354,855 (specifically incorporated herein by reference) reports that certain ribozymes can act as endonucleases with a sequence
10 specificity greater than that of known ribonucleases and approaching that of the DNA restriction enzymes. Thus, sequence-specific ribozyme-mediated inhibition of gene expression may be particularly suited to therapeutic applications (Scanlon *et al.*, 1991; Sarver *et al.*, 1990). Recently, it was reported that ribozymes elicited genetic changes in some cells lines to which they were applied; the altered genes included the oncogenes
15 H-ras, c-fos and genes of HIV. Most of this work involved the modification of a target mRNA, based on a specific mutant codon that is cleaved by a specific ribozyme.

Six basic varieties of naturally-occurring enzymatic RNAs are known presently. Each can catalyze the hydrolysis of RNA phosphodiester bonds *in trans* (and thus can cleave other RNA molecules) under physiological conditions. In general,
20 enzymatic nucleic acids act by first binding to a target RNA. Such binding occurs through the target binding portion of a enzymatic nucleic acid which is held in close proximity to an enzymatic portion of the molecule that acts to cleave the target RNA. Thus, the enzymatic nucleic acid first recognizes and then binds a target RNA through complementary base-pairing, and once bound to the correct site, acts enzymatically to
25 cut the target RNA. Strategic cleavage of such a target RNA will destroy its ability to direct synthesis of an encoded protein. After an enzymatic nucleic acid has bound and cleaved its RNA target, it is released from that RNA to search for another target and can repeatedly bind and cleave new targets.

The enzymatic nature of a ribozyme is advantageous over many
30 technologies, such as antisense technology (where a nucleic acid molecule simply binds to a nucleic acid target to block its translation) since the concentration of ribozyme

necessary to affect a therapeutic treatment is lower than that of an antisense oligonucleotide. This advantage reflects the ability of the ribozyme to act enzymatically. Thus, a single ribozyme molecule is able to cleave many molecules of target RNA. In addition, the ribozyme is a highly specific inhibitor, with the specificity 5 of inhibition depending not only on the base pairing mechanism of binding to the target RNA, but also on the mechanism of target RNA cleavage. Single mismatches, or base-substitutions, near the site of cleavage can completely eliminate catalytic activity of a ribozyme. Similar mismatches in antisense molecules do not prevent their action (Woolf *et al.*, 1992). Thus, the specificity of action of a ribozyme is greater than that of 10 an antisense oligonucleotide binding the same RNA site.

The enzymatic nucleic acid molecule may be formed in a hammerhead, hairpin, a hepatitis δ virus, group I intron or RNaseP RNA (in association with an RNA guide sequence) or Neurospora VS RNA motif. Examples of hammerhead motifs are described by Rossi *et al.* (1992). Examples of hairpin motifs are described by Hampel 15 *et al.* (Eur. Pat. Appl. Publ. No. EP 0360257), Hampel and Tritz (1989), Hampel *et al.* (1990) and U. S. Patent 5,631,359 (specifically incorporated herein by reference). An example of the hepatitis δ virus motif is described by Perrotta and Been (1992); an example of the RNaseP motif is described by Guerrier-Takada *et al.* (1983); Neurospora 20 VS RNA ribozyme motif is described by Collins (Saville and Collins, 1990; Saville and Collins, 1991; Collins and Olive, 1993); and an example of the Group I intron is described in (U. S. Patent 4,987,071, specifically incorporated herein by reference). All that is important in an enzymatic nucleic acid molecule of this invention is that it has a specific substrate binding site which is complementary to one or more of the target gene 25 RNA regions, and that it have nucleotide sequences within or surrounding that substrate binding site which impart an RNA cleaving activity to the molecule. Thus the ribozyme constructs need not be limited to specific motifs mentioned herein.

In certain embodiments, it may be important to produce enzymatic cleaving agents which exhibit a high degree of specificity for the RNA of a desired target, such as one of the sequences disclosed herein. The enzymatic nucleic acid 30 molecule is preferably targeted to a highly conserved sequence region of a target mRNA. Such enzymatic nucleic acid molecules can be delivered exogenously to

specific cells as required. Alternatively, the ribozymes can be expressed from DNA or RNA vectors that are delivered to specific cells.

Small enzymatic nucleic acid motifs (*e.g.*, of the hammerhead or the hairpin structure) may also be used for exogenous delivery. The simple structure of 5 these molecules increases the ability of the enzymatic nucleic acid to invade targeted regions of the mRNA structure. Alternatively, catalytic RNA molecules can be expressed within cells from eukaryotic promoters (*e.g.*, Scanlon *et al.*, 1991; Kashani-Sabet *et al.*, 1992; Dropulic *et al.*, 1992; Weerasinghe *et al.*, 1991; Ojwang *et al.*, 1992; Chen *et al.*, 1992; Sarver *et al.*, 1990). Those skilled in the art realize that any ribozyme 10 can be expressed in eukaryotic cells from the appropriate DNA vector. The activity of such ribozymes can be augmented by their release from the primary transcript by a second ribozyme (Int. Pat. Appl. Publ. No. WO 93/23569, and Int. Pat. Appl. Publ. No. WO 94/02595, both hereby incorporated by reference; Ohkawa *et al.*, 1992; Taira *et al.*, 1991; and Ventura *et al.*, 1993).

15 Ribozymes may be added directly, or can be complexed with cationic lipids, lipid complexes, packaged within liposomes, or otherwise delivered to target cells. The RNA or RNA complexes can be locally administered to relevant tissues *ex vivo*, or *in vivo* through injection, aerosol inhalation, infusion pump or stent, with or without their incorporation in biopolymers.

20 Ribozymes may be designed as described in Int. Pat. Appl. Publ. No. WO 93/23569 and Int. Pat. Appl. Publ. No. WO 94/02595, each specifically incorporated herein by reference) and synthesized to be tested *in vitro* and *in vivo*, as described. Such ribozymes can also be optimized for delivery. While specific examples are provided, those in the art will recognize that equivalent RNA targets in 25 other species can be utilized when necessary.

Hammerhead or hairpin ribozymes may be individually analyzed by computer folding (Jaeger *et al.*, 1989) to assess whether the ribozyme sequences fold into the appropriate secondary structure. Those ribozymes with unfavorable intramolecular interactions between the binding arms and the catalytic core are 30 eliminated from consideration. Varying binding arm lengths can be chosen to optimize

activity. Generally, at least 5 or so bases on each arm are able to bind to, or otherwise interact with, the target RNA.

Ribozymes of the hammerhead or hairpin motif may be designed to anneal to various sites in the mRNA message, and can be chemically synthesized. The 5 method of synthesis used follows the procedure for normal RNA synthesis as described in Usman *et al.* (1987) and in Scaringe *et al.* (1990) and makes use of common nucleic acid protecting and coupling groups, such as dimethoxytrityl at the 5'-end, and phosphoramidites at the 3'-end. Average stepwise coupling yields are typically >98%. Hairpin ribozymes may be synthesized in two parts and annealed to reconstruct an 10 active ribozyme (Chowrira and Burke, 1992). Ribozymes may be modified extensively to enhance stability by modification with nuclease resistant groups, for example, 2'-amino, 2'-C-allyl, 2'-flouro, 2'-o-methyl, 2'-H (for a review see e.g., Usman and Cedergren, 1992). Ribozymes may be purified by gel electrophoresis using general methods or by high pressure liquid chromatography and resuspended in water.

15 Ribozyme activity can be optimized by altering the length of the ribozyme binding arms, or chemically synthesizing ribozymes with modifications that prevent their degradation by serum ribonucleases (see e.g., Int. Pat. Appl. Publ. No. WO 92/07065; Perrault *et al.*, 1990; Pieken *et al.*, 1991; Usman and Cedergren, 1992; Int. Pat. Appl. Publ. No. WO 93/15187; Int. Pat. Appl. Publ. No. WO 91/03162; Eur. Pat. 20 Appl. Publ. No. 92110298.4; U. S. Patent 5,334,711; and Int. Pat. Appl. Publ. No. WO 94/13688, which describe various chemical modifications that can be made to the sugar moieties of enzymatic RNA molecules), modifications which enhance their efficacy in cells, and removal of stem II bases to shorten RNA synthesis times and reduce chemical requirements.

25 Sullivan *et al.* (Int. Pat. Appl. Publ. No. WO 94/02595) describes the general methods for delivery of enzymatic RNA molecules. Ribozymes may be administered to cells by a variety of methods known to those familiar to the art, including, but not restricted to, encapsulation in liposomes, by iontophoresis, or by incorporation into other vehicles, such as hydrogels, cyclodextrins, biodegradable 30 nanocapsules, and bioadhesive microspheres. For some indications, ribozymes may be directly delivered *ex vivo* to cells or tissues with or without the aforementioned vehicles.

Alternatively, the RNA/vehicle combination may be locally delivered by direct inhalation, by direct injection or by use of a catheter, infusion pump or stent. Other routes of delivery include, but are not limited to, intravascular, intramuscular, subcutaneous or joint injection, aerosol inhalation, oral (tablet or pill form), topical, 5 systemic, ocular, intraperitoneal and/or intrathecal delivery. More detailed descriptions of ribozyme delivery and administration are provided in Int. Pat. Appl. Publ. No. WO 94/02595 and Int. Pat. Appl. Publ. No. WO 93/23569, each specifically incorporated herein by reference.

Another means of accumulating high concentrations of a ribozyme(s) 10 within cells is to incorporate the ribozyme-encoding sequences into a DNA expression vector. Transcription of the ribozyme sequences are driven from a promoter for eukaryotic RNA polymerase I (pol I), RNA polymerase II (pol II), or RNA polymerase III (pol III). Transcripts from pol II or pol III promoters will be expressed at high levels in all cells; the levels of a given pol II promoter in a given cell type will depend on the 15 nature of the gene regulatory sequences (enhancers, silencers, *etc.*) present nearby. Prokaryotic RNA polymerase promoters may also be used, providing that the prokaryotic RNA polymerase enzyme is expressed in the appropriate cells (Elroy-Stein and Moss, 1990; Gao and Huang, 1993; Lieber *et al.*, 1993; Zhou *et al.*, 1990). Ribozymes expressed from such promoters can function in mammalian cells (*e.g.* 20 Kashani-Saber *et al.*, 1992; Ojwang *et al.*, 1992; Chen *et al.*, 1992; Yu *et al.*, 1993; L'Huillier *et al.*, 1992; Lisziewicz *et al.*, 1993). Such transcription units can be incorporated into a variety of vectors for introduction into mammalian cells, including but not restricted to, plasmid DNA vectors, viral DNA vectors (such as adenovirus or 25 adeno-associated vectors), or viral RNA vectors (such as retroviral, semliki forest virus, sindbis virus vectors).

Ribozymes may be used as diagnostic tools to examine genetic drift and mutations within diseased cells. They can also be used to assess levels of the target RNA molecule. The close relationship between ribozyme activity and the structure of the target RNA allows the detection of mutations in any region of the molecule which 30 alters the base-pairing and three-dimensional structure of the target RNA. By using multiple ribozymes, one may map nucleotide changes which are important to RNA

structure and function *in vitro*, as well as in cells and tissues. Cleavage of target RNAs with ribozymes may be used to inhibit gene expression and define the role (essentially) of specified gene products in the progression of disease. In this manner, other genetic targets may be defined as important mediators of the disease. These studies will lead to 5 better treatment of the disease progression by affording the possibility of combinational therapies (e.g., multiple ribozymes targeted to different genes, ribozymes coupled with known small molecule inhibitors, or intermittent treatment with combinations of ribozymes and/or other chemical or biological molecules). Other *in vitro* uses of ribozymes are well known in the art, and include detection of the presence of mRNA 10 associated with an IL-5 related condition. Such RNA is detected by determining the presence of a cleavage product after treatment with a ribozyme using standard methodology.

PEPTIDE NUCLEIC ACIDS

In certain embodiments, the inventors contemplate the use of peptide 15 nucleic acids (PNAs) in the practice of the methods of the invention. PNA is a DNA mimic in which the nucleobases are attached to a pseudopeptide backbone (Good and Nielsen, 1997). PNA is able to be utilized in a number methods that traditionally have used RNA or DNA. Often PNA sequences perform better in techniques than the corresponding RNA or DNA sequences and have utilities that are not inherent to RNA 20 or DNA. A review of PNA including methods of making, characteristics of, and methods of using, is provided by Corey (1997) and is incorporated herein by reference. As such, in certain embodiments, one may prepare PNA sequences that are complementary to one or more portions of the ACE mRNA sequence, and such PNA compositions may be used to regulate, alter, decrease, or reduce the translation of ACE- 25 specific mRNA, and thereby alter the level of ACE activity in a host cell to which such PNA compositions have been administered.

PNAs have 2-aminoethyl-glycine linkages replacing the normal phosphodiester backbone of DNA (Nielsen *et al.*, 1991; Hanvey *et al.*, 1992; Hyrup and Nielsen, 1996; Neilsen, 1996). This chemistry has three important consequences: 30 firstly, in contrast to DNA or phosphorothioate oligonucleotides, PNAs are neutral

molecules; secondly, PNAs are achiral, which avoids the need to develop a stereoselective synthesis; and thirdly, PNA synthesis uses standard Boc (Dueholm *et al.*, 1994) or Fmoc (Thomson *et al.*, 1995) protocols for solid-phase peptide synthesis, although other methods, including a modified Merrifield method, have been used
5 (Christensen *et al.*, 1995).

PNA monomers or ready-made oligomers are commercially available from PerSeptive Biosystems (Framingham, MA). PNA syntheses by either Boc or Fmoc protocols are straightforward using manual or automated protocols (Norton *et al.*, 1995). The manual protocol lends itself to the production of chemically modified PNAs
10 or the simultaneous synthesis of families of closely related PNAs.

As with peptide synthesis, the success of a particular PNA synthesis will depend on the properties of the chosen sequence. For example, while in theory PNAs can incorporate any combination of nucleotide bases, the presence of adjacent purines can lead to deletions of one or more residues in the product. In expectation of this
15 difficulty, it is suggested that, in producing PNAs with adjacent purines, one should repeat the coupling of residues likely to be added inefficiently. This should be followed by the purification of PNAs by reverse-phase high-pressure liquid chromatography (Norton *et al.*, 1995) providing yields and purity of product similar to those observed during the synthesis of peptides.

Modifications of PNAs for a given application may be accomplished by coupling amino acids during solid-phase synthesis or by attaching compounds that contain a carboxylic acid group to the exposed N-terminal amine. Alternatively, PNAs can be modified after synthesis by coupling to an introduced lysine or cysteine. The ease with which PNAs can be modified facilitates optimization for better solubility or
20 for specific functional requirements. Once synthesized, the identity of PNAs and their derivatives can be confirmed by mass spectrometry. Several studies have made and utilized modifications of PNAs (Norton *et al.*, 1995; Haaima *et al.*, 1996; Stetsenko *et al.*, 1996; Petersen *et al.*, 1995; Ulmann *et al.*, 1996; Koch *et al.*, 1995; Orum *et al.*, 1995; Footer *et al.*, 1996; Griffith *et al.*, 1995; Kremsky *et al.*, 1996; Pardridge *et al.*,
25 1995; Boffa *et al.*, 1995; Landsdorp *et al.*, 1996; Gambacorti-Passerini *et al.*, 1996;
30 Armitage *et al.*, 1997; Seeger *et al.*, 1997; Ruskowski *et al.*, 1997). U.S. Patent No.

5,700,922 discusses PNA-DNA-PNA chimeric molecules and their uses in diagnostics, modulating protein in organisms, and treatment of conditions susceptible to therapeutics.

In contrast to DNA and RNA, which contain negatively charged linkages, the PNA backbone is neutral. In spite of this dramatic alteration, PNAs recognize complementary DNA and RNA by Watson-Crick pairing (Egholm *et al.*, 1993), validating the initial modeling by Nielsen *et al.* (1991). PNAs lack 3' to 5' polarity and can bind in either parallel or antiparallel fashion, with the antiparallel mode being preferred (Egholm *et al.*, 1993).

10 Hybridization of DNA oligonucleotides to DNA and RNA is destabilized by electrostatic repulsion between the negatively charged phosphate backbones of the complementary strands. By contrast, the absence of charge repulsion in PNA-DNA or PNA-RNA duplexes increases the melting temperature (T_m) and reduces the dependence of T_m on the concentration of mono- or divalent cations (Nielsen *et al.*, 1991). The
15 enhanced rate and affinity of hybridization are significant because they are responsible for the surprising ability of PNAs to perform strand invasion of complementary sequences within relaxed double-stranded DNA. In addition, the efficient hybridization at inverted repeats suggests that PNAs can recognize secondary structure effectively within double-stranded DNA. Enhanced recognition also occurs with PNAs
20 immobilized on surfaces, and Wang *et al.* have shown that support-bound PNAs can be used to detect hybridization events (Wang *et al.*, 1996).

One might expect that tight binding of PNAs to complementary sequences would also increase binding to similar (but not identical) sequences, reducing the sequence specificity of PNA recognition. As with DNA hybridization, however,
25 selective recognition can be achieved by balancing oligomer length and incubation temperature. Moreover, selective hybridization of PNAs is encouraged by PNA-DNA hybridization being less tolerant of base mismatches than DNA-DNA hybridization. For example, a single mismatch within a 16 bp PNA-DNA duplex can reduce the T_m by up to 15°C (Egholm *et al.*, 1993). This high level of discrimination has allowed the
30 development of several PNA-based strategies for the analysis of point mutations (Wang

et al., 1996; Carlsson *et al.*, 1996; Thiede *et al.*, 1996; Webb and Hurskainen, 1996; Perry-O'Keefe *et al.*, 1996).

High-affinity binding provides clear advantages for molecular recognition and the development of new applications for PNAs. For example, 11-13 5 nucleotide PNAs inhibit the activity of telomerase, a ribonucleo-protein that extends telomere ends using an essential RNA template, while the analogous DNA oligomers do not (Norton *et al.*, 1996).

Neutral PNAs are more hydrophobic than analogous DNA oligomers, and this can lead to difficulty solubilizing them at neutral pH, especially if the PNAs 10 have a high purine content or if they have the potential to form secondary structures. Their solubility can be enhanced by attaching one or more positive charges to the PNA termini (Nielsen *et al.*, 1991).

Findings by Allfrey and colleagues suggest that strand invasion will occur spontaneously at sequences within chromosomal DNA (Boffa *et al.*, 1995; Boffa 15 *et al.*, 1996). These studies targeted PNAs to triplet repeats of the nucleotides CAG and used this recognition to purify transcriptionally active DNA (Boffa *et al.*, 1995) and to inhibit transcription (Boffa *et al.*, 1996). This result suggests that if PNAs can be delivered within cells then they will have the potential to be general sequence-specific regulators of gene expression. Studies and reviews concerning the use of PNAs as 20 antisense and anti-gene agents include Nielsen *et al.* (1993b), Hanvey *et al.* (1992), and Good and Nielsen (1997). Koppelhus *et al.* (1997) have used PNAs to inhibit HIV-1 inverse transcription, showing that PNAs may be used for antiviral therapies.

Methods of characterizing the antisense binding properties of PNAs are discussed in Rose (1993) and Jensen *et al.* (1997). Rose uses capillary gel 25 electrophoresis to determine binding of PNAs to their complementary oligonucleotide, measuring the relative binding kinetics and stoichiometry. Similar types of measurements were made by Jensen *et al.* using BIACore™ technology.

Other applications of PNAs include use in DNA strand invasion (Nielsen *et al.*, 1991), antisense inhibition (Hanvey *et al.*, 1992), mutational analysis (Orum *et 30 al.*, 1993), enhancers of transcription (Mollegaard *et al.*, 1994), nucleic acid purification (Orum *et al.*, 1995), isolation of transcriptionally active genes (Boffa *et al.*, 1995),

blocking of transcription factor binding (Vickers *et al.*, 1995), genome cleavage (Veselkov *et al.*, 1996), biosensors (Wang *et al.*, 1996), *in situ* hybridization (Thisted *et al.*, 1996), and in an alternative to Southern blotting (Perry-O'Keefe, 1996).

POLYPEPTIDE COMPOSITIONS

5 The present invention, in other aspects, provides polypeptide compositions. Generally, a polypeptide of the invention will be an isolated polypeptide (or an epitope, variant, or active fragment thereof) derived from a mammalian species. Preferably, the polypeptide is encoded by a polynucleotide sequence disclosed herein or 10 a sequence which hybridizes under moderately stringent conditions to a polynucleotide sequence disclosed herein. Alternatively, the polypeptide may be defined as a polypeptide which comprises a contiguous amino acid sequence from an amino acid sequence disclosed herein, or which polypeptide comprises an entire amino acid sequence disclosed herein.

In the present invention, a polypeptide composition is also understood to 15 comprise one or more polypeptides that are immunologically reactive with antibodies generated against a polypeptide of the invention, or to active fragments, or to variants or biological functional equivalents thereof.

Likewise, a polypeptide composition of the present invention is understood to comprise one or more polypeptides that are capable of eliciting antibodies 20 that are immunologically reactive with one or more polypeptides encoded by one or more contiguous nucleic acid sequences contained in SEQ ID NO: 1-38, 42-204, 205, 207 and 210-290, or to active fragments, or to variants thereof, or to one or more nucleic acid sequences which hybridize to one or more of these sequences under conditions of moderate to high stringency.

25 As used herein, an active fragment of a polypeptide includes a whole or a portion of a polypeptide which is modified by conventional techniques, *e.g.*, mutagenesis, or by addition, deletion, or substitution, but which active fragment exhibits substantially the same structure function, antigenicity, etc., as a polypeptide as described herein.

In certain illustrative embodiments, the polypeptides of the invention will comprise at least an immunogenic portion of a breast tumor protein or a variant thereof, as described herein. As noted above, a "breast tumor protein" is a protein that is expressed by breast tumor cells. Proteins that are breast tumor proteins react 5 detectably within an immunoassay (such as an ELISA) with antisera from a patient with breast cancer. Polypeptides as described herein may be of any length. Additional sequences derived from the native protein and/or heterologous sequences may be present, and such sequences may (but need not) possess further immunogenic or antigenic properties.

10 An "immunogenic portion," as used herein is a portion of a protein that is recognized (*i.e.*, specifically bound) by a B-cell and/or T-cell surface antigen receptor. Such immunogenic portions generally comprise at least 5 amino acid residues, more preferably at least 10, and still more preferably at least 20 amino acid residues of a breast tumor protein or a variant thereof. Certain preferred immunogenic 15 portions include peptides in which an N-terminal leader sequence and/or transmembrane domain have been deleted. Other preferred immunogenic portions may contain a small N- and/or C-terminal deletion (*e.g.*, 1-30 amino acids, preferably 5-15 amino acids), relative to the mature protein.

Immunogenic portions may generally be identified using well known 20 techniques, such as those summarized in Paul, *Fundamental Immunology*, 3rd ed., 243-247 (Raven Press, 1993) and references cited therein. Such techniques include screening polypeptides for the ability to react with antigen-specific antibodies, antisera and/or T-cell lines or clones. As used herein, antisera and antibodies are "antigen-specific" if they specifically bind to an antigen (*i.e.*, they react with the protein in an 25 ELISA or other immunoassay, and do not react detectably with unrelated proteins). Such antisera and antibodies may be prepared as described herein, and using well known techniques. An immunogenic portion of a native breast tumor protein is a portion that reacts with such antisera and/or T-cells at a level that is not substantially less than the reactivity of the full length polypeptide (*e.g.*, in an ELISA and/or T-cell 30 reactivity assay). Such immunogenic portions may react within such assays at a level that is similar to or greater than the reactivity of the full length polypeptide. Such

screens may generally be performed using methods well known to those of ordinary skill in the art, such as those described in Harlow and Lane, *Antibodies: A Laboratory Manual*, Cold Spring Harbor Laboratory, 1988. For example, a polypeptide may be immobilized on a solid support and contacted with patient sera to allow binding of 5 antibodies within the sera to the immobilized polypeptide. Unbound sera may then be removed and bound antibodies detected using, for example, ¹²⁵I-labeled Protein A.

As noted above, a composition may comprise a variant of a native breast tumor protein. A polypeptide "variant," as used herein, is a polypeptide that differs from a native breast tumor protein in one or more substitutions, deletions, additions 10 and/or insertions, such that the immunogenicity of the polypeptide is not substantially diminished. In other words, the ability of a variant to react with antigen-specific antisera may be enhanced or unchanged, relative to the native protein, or may be diminished by less than 50%, and preferably less than 20%, relative to the native protein. Such variants may generally be identified by modifying one of the above 15 polypeptide sequences and evaluating the reactivity of the modified polypeptide with antigen-specific antibodies or antisera as described herein. Preferred variants include those in which one or more portions, such as an N-terminal leader sequence or transmembrane domain, have been removed. Other preferred variants include variants in which a small portion (e.g., 1-30 amino acids, preferably 5-15 amino acids) has been 20 removed from the N- and/or C-terminal of the mature protein.

Polypeptide variants encompassed by the present invention include those exhibiting at least about 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, or 99% or more identity (determined as described above) to the polypeptides disclosed herein.

25 Preferably, a variant contains conservative substitutions. A "conservative substitution" is one in which an amino acid is substituted for another amino acid that has similar properties, such that one skilled in the art of peptide chemistry would expect the secondary structure and hydrophobic nature of the polypeptide to be substantially unchanged. Amino acid substitutions may generally be 30 made on the basis of similarity in polarity, charge, solubility, hydrophobicity, hydrophilicity and/or the amphipathic nature of the residues. For example, negatively

charged amino acids include aspartic acid and glutamic acid; positively charged amino acids include lysine and arginine; and amino acids with uncharged polar head groups having similar hydrophilicity values include leucine, isoleucine and valine; glycine and alanine; asparagine and glutamine; and serine, threonine, phenylalanine and tyrosine.

- 5 Other groups of amino acids that may represent conservative changes include: (1) ala, pro, gly, glu, asp, gln, asn, ser, thr; (2) cys, ser, tyr, thr; (3) val, ile, leu, met, ala, phe; (4) lys, arg, his; and (5) phe, tyr, trp, his. A variant may also, or alternatively, contain nonconservative changes. In a preferred embodiment, variant polypeptides differ from a native sequence by substitution, deletion or addition of five amino acids or fewer.
- 10 Variants may also (or alternatively) be modified by, for example, the deletion or addition of amino acids that have minimal influence on the immunogenicity, secondary structure and hydropathic nature of the polypeptide.

As noted above, polypeptides may comprise a signal (or leader) sequence at the N-terminal end of the protein, which co-translationally or post-translationally directs transfer of the protein. The polypeptide may also be conjugated to a linker or other sequence for ease of synthesis, purification or identification of the polypeptide (e.g., poly-His), or to enhance binding of the polypeptide to a solid support. For example, a polypeptide may be conjugated to an immunoglobulin Fc region.

Polypeptides may be prepared using any of a variety of well known techniques. Recombinant polypeptides encoded by DNA sequences as described above may be readily prepared from the DNA sequences using any of a variety of expression vectors known to those of ordinary skill in the art. Expression may be achieved in any appropriate host cell that has been transformed or transfected with an expression vector containing a DNA molecule that encodes a recombinant polypeptide. Suitable host cells include prokaryotes, yeast, and higher eukaryotic cells, such as mammalian cells and plant cells. Preferably, the host cells employed are *E. coli*, yeast or a mammalian cell line such as COS or CHO. Supernatants from suitable host/vector systems which secrete recombinant protein or polypeptide into culture media may be first concentrated using a commercially available filter. Following concentration, the concentrate may be applied to a suitable purification matrix such as an affinity matrix or an ion exchange

resin. Finally, one or more reverse phase HPLC steps can be employed to further purify a recombinant polypeptide.

Portions and other variants having less than about 100 amino acids, and generally less than about 50 amino acids, may also be generated by synthetic means, 5 using techniques well known to those of ordinary skill in the art. For example, such polypeptides may be synthesized using any of the commercially available solid-phase techniques, such as the Merrifield solid-phase synthesis method, where amino acids are sequentially added to a growing amino acid chain. See Merrifield, *J. Am. Chem. Soc.* 85:2149-2146, 1963. Equipment for automated synthesis of polypeptides is 10 commercially available from suppliers such as Perkin Elmer/Applied BioSystems Division (Foster City, CA), and may be operated according to the manufacturer's instructions.

Within certain specific embodiments, a polypeptide may be a fusion protein that comprises multiple polypeptides as described herein, or that comprises at 15 least one polypeptide as described herein and an unrelated sequence, such as a known tumor protein. A fusion partner may, for example, assist in providing T helper epitopes (an immunological fusion partner), preferably T helper epitopes recognized by humans, or may assist in expressing the protein (an expression enhancer) at higher yields than the native recombinant protein. Certain preferred fusion partners are both 20 immunological and expression enhancing fusion partners. Other fusion partners may be selected so as to increase the solubility of the protein or to enable the protein to be targeted to desired intracellular compartments. Still further fusion partners include affinity tags, which facilitate purification of the protein.

Fusion proteins may generally be prepared using standard techniques, 25 including chemical conjugation. Preferably, a fusion protein is expressed as a recombinant protein, allowing the production of increased levels, relative to a non-fused protein, in an expression system. Briefly, DNA sequences encoding the polypeptide components may be assembled separately, and ligated into an appropriate expression vector. The 3' end of the DNA sequence encoding one polypeptide component is 30 ligated, with or without a peptide linker, to the 5' end of a DNA sequence encoding the second polypeptide component so that the reading frames of the sequences are in phase.

This permits translation into a single fusion protein that retains the biological activity of both component polypeptides.

A peptide linker sequence may be employed to separate the first and second polypeptide components by a distance sufficient to ensure that each polypeptide 5 folds into its secondary and tertiary structures. Such a peptide linker sequence is incorporated into the fusion protein using standard techniques well known in the art. Suitable peptide linker sequences may be chosen based on the following factors: (1) their ability to adopt a flexible extended conformation; (2) their inability to adopt a secondary structure that could interact with functional epitopes on the first and second 10 polypeptides; and (3) the lack of hydrophobic or charged residues that might react with the polypeptide functional epitopes. Preferred peptide linker sequences contain Gly, Asn and Ser residues. Other near neutral amino acids, such as Thr and Ala may also be used in the linker sequence. Amino acid sequences which may be usefully employed as linkers include those disclosed in Maratea et al., *Gene* 40:39-46, 1985; Murphy et al., 15 *Proc. Natl. Acad. Sci. USA* 83:8258-8262, 1986; U.S. Patent No. 4,935,233 and U.S. Patent No. 4,751,180. The linker sequence may generally be from 1 to about 50 amino acids in length. Linker sequences are not required when the first and second polypeptides have non-essential N-terminal amino acid regions that can be used to separate the functional domains and prevent steric interference.

20 The ligated DNA sequences are operably linked to suitable transcriptional or translational regulatory elements. The regulatory elements responsible for expression of DNA are located only 5' to the DNA sequence encoding the first polypeptides. Similarly, stop codons required to end translation and transcription termination signals are only present 3' to the DNA sequence encoding the 25 second polypeptide.

Fusion proteins are also provided. Such proteins comprise a polypeptide as described herein together with an unrelated immunogenic protein. Preferably the immunogenic protein is capable of eliciting a recall response. Examples of such proteins include tetanus, tuberculosis and hepatitis proteins (*see, for example, Stoute 30 et al. New Engl. J. Med.*, 336:86-91, 1997).

Within preferred embodiments, an immunological fusion partner is derived from protein D, a surface protein of the gram-negative bacterium Haemophilus influenza B (WO 91/18926). Preferably, a protein D derivative comprises approximately the first third of the protein (e.g., the first N-terminal 100-110 amino acids), and a protein D derivative may be lipidated. Within certain preferred embodiments, the first 109 residues of a Lipoprotein D fusion partner is included on the N-terminus to provide the polypeptide with additional exogenous T-cell epitopes and to increase the expression level in *E. coli* (thus functioning as an expression enhancer). The lipid tail ensures optimal presentation of the antigen to antigen presenting cells.

10 Other fusion partners include the non-structural protein from influenzae virus, NS1 (hemagglutinin). Typically, the N-terminal 81 amino acids are used, although different fragments that include T-helper epitopes may be used.

In another embodiment, the immunological fusion partner is the protein known as LYTA, or a portion thereof (preferably a C-terminal portion). LYTA is derived from *Streptococcus pneumoniae*, which synthesizes an N-acetyl-L-alanine amidase known as amidase LYTA (encoded by the LytA gene; *Gene* 43:265-292, 1986). LYTA is an autolysin that specifically degrades certain bonds in the peptidoglycan backbone. The C-terminal domain of the LYTA protein is responsible for the affinity to the choline or to some choline analogues such as DEAE. This property has been exploited for the development of *E. coli* C-LYTA expressing plasmids useful for expression of fusion proteins. Purification of hybrid proteins containing the C-LYTA fragment at the amino terminus has been described (see *Biotechnology* 10:795-798, 1992). Within a preferred embodiment, a repeat portion of LYTA may be incorporated into a fusion protein. A repeat portion is found in the C-terminal region starting at residue 178. A particularly preferred repeat portion incorporates residues 188-305.

In general, polypeptides (including fusion proteins) and polynucleotides as described herein are isolated. An "isolated" polypeptide or polynucleotide is one that is removed from its original environment. For example, a naturally-occurring protein is isolated if it is separated from some or all of the coexisting materials in the natural system. Preferably, such polypeptides are at least about 90% pure, more preferably at

least about 95% pure and most preferably at least about 99% pure. A polynucleotide is considered to be isolated if, for example, it is cloned into a vector that is not a part of the natural environment.

BINDING AGENTS

5 The present invention further provides agents, such as antibodies and antigen-binding fragments thereof, that specifically bind to a breast tumor protein. As used herein, an antibody, or antigen-binding fragment thereof, is said to "specifically bind" to a breast tumor protein if it reacts at a detectable level (within, for example, an ELISA) with a breast tumor protein, and does not react detectably with unrelated 10 proteins under similar conditions. As used herein, "binding" refers to a noncovalent association between two separate molecules such that a complex is formed. The ability to bind may be evaluated by, for example, determining a binding constant for the formation of the complex. The binding constant is the value obtained when the concentration of the complex is divided by the product of the component 15 concentrations. In general, two compounds are said to "bind," in the context of the present invention, when the binding constant for complex formation exceeds about 10^3 L/mol. The binding constant may be determined using methods well known in the art.

Binding agents may be further capable of differentiating between patients with and without a cancer, such as breast cancer, using the representative assays 20 provided herein. In other words, antibodies or other binding agents that bind to a breast tumor protein will generate a signal indicating the presence of a cancer in at least about 20% of patients with the disease, and will generate a negative signal indicating the absence of the disease in at least about 90% of individuals without the cancer. To determine whether a binding agent satisfies this requirement, biological samples (*e.g.*, 25 blood, sera, sputum, urine and/or tumor biopsies) from patients with and without a cancer (as determined using standard clinical tests) may be assayed as described herein for the presence of polypeptides that bind to the binding agent. It will be apparent that a statistically significant number of samples with and without the disease should be assayed. Each binding agent should satisfy the above criteria; however, those of

ordinary skill in the art will recognize that binding agents may be used in combination to improve sensitivity.

Any agent that satisfies the above requirements may be a binding agent. For example, a binding agent may be a ribosome, with or without a peptide component, 5 an RNA molecule or a polypeptide. In a preferred embodiment, a binding agent is an antibody or an antigen-binding fragment thereof. Antibodies may be prepared by any of a variety of techniques known to those of ordinary skill in the art. *See, e.g.*, Harlow and Lane, *Antibodies: A Laboratory Manual*, Cold Spring Harbor Laboratory, 1988. In general, antibodies can be produced by cell culture techniques, including the generation 10 of monoclonal antibodies as described herein, or via transfection of antibody genes into suitable bacterial or mammalian cell hosts, in order to allow for the production of recombinant antibodies. In one technique, an immunogen comprising the polypeptide is initially injected into any of a wide variety of mammals (*e.g.*, mice, rats, rabbits, sheep or goats). In this step, the polypeptides of this invention may serve as the immunogen 15 without modification. Alternatively, particularly for relatively short polypeptides, a superior immune response may be elicited if the polypeptide is joined to a carrier protein, such as bovine serum albumin or keyhole limpet hemocyanin. The immunogen is injected into the animal host, preferably according to a predetermined schedule incorporating one or more booster immunizations, and the animals are bled periodically. 20 Polyclonal antibodies specific for the polypeptide may then be purified from such antisera by, for example, affinity chromatography using the polypeptide coupled to a suitable solid support.

Monoclonal antibodies specific for an antigenic polypeptide of interest may be prepared, for example, using the technique of Kohler and Milstein, *Eur. J. 25 Immunol.* 6:511-519, 1976, and improvements thereto. Briefly, these methods involve the preparation of immortal cell lines capable of producing antibodies having the desired specificity (*i.e.*, reactivity with the polypeptide of interest). Such cell lines may be produced, for example, from spleen cells obtained from an animal immunized as described above. The spleen cells are then immortalized by, for example, fusion with a 30 myeloma cell fusion partner, preferably one that is syngeneic with the immunized animal. A variety of fusion techniques may be employed. For example, the spleen cells

and myeloma cells may be combined with a nonionic detergent for a few minutes and then plated at low density on a selective medium that supports the growth of hybrid cells, but not myeloma cells. A preferred selection technique uses HAT (hypoxanthine, aminopterin, thymidine) selection. After a sufficient time, usually about 1 to 2 weeks, 5 colonies of hybrids are observed. Single colonies are selected and their culture supernatants tested for binding activity against the polypeptide. Hybridomas having high reactivity and specificity are preferred.

Monoclonal antibodies may be isolated from the supernatants of growing hybridoma colonies. In addition, various techniques may be employed to enhance the 10 yield, such as injection of the hybridoma cell line into the peritoneal cavity of a suitable vertebrate host, such as a mouse. Monoclonal antibodies may then be harvested from the ascites fluid or the blood. Contaminants may be removed from the antibodies by conventional techniques, such as chromatography, gel filtration, precipitation, and extraction. The polypeptides of this invention may be used in the purification process 15 in, for example, an affinity chromatography step.

Within certain embodiments, the use of antigen-binding fragments of antibodies may be preferred. Such fragments include Fab fragments, which may be prepared using standard techniques. Briefly, immunoglobulins may be purified from rabbit serum by affinity chromatography on Protein A bead columns (Harlow and Lane, 20 *Antibodies: A Laboratory Manual*, Cold Spring Harbor Laboratory, 1988) and digested by papain to yield Fab and Fc fragments. The Fab and Fc fragments may be separated by affinity chromatography on protein A bead columns.

Monoclonal antibodies of the present invention may be coupled to one or more therapeutic agents. Suitable agents in this regard include radionuclides, 25 differentiation inducers, drugs, toxins, and derivatives thereof. Preferred radionuclides include ⁹⁰Y, ¹²³I, ¹²⁵I, ¹³¹I, ¹⁸⁶Re, ¹⁸⁸Re, ²¹¹At, and ²¹²Bi. Preferred drugs include methotrexate, and pyrimidine and purine analogs. Preferred differentiation inducers include phorbol esters and butyric acid. Preferred toxins include ricin, abrin, diphteria toxin, cholera toxin, gelonin, *Pseudomonas* exotoxin, *Shigella* toxin, and pokeweed 30 antiviral protein.

A therapeutic agent may be coupled (*e.g.*, covalently bonded) to a suitable monoclonal antibody either directly or indirectly (*e.g.*, via a linker group). A direct reaction between an agent and an antibody is possible when each possesses a substituent capable of reacting with the other. For example, a nucleophilic group, such 5 as an amino or sulphydryl group, on one may be capable of reacting with a carbonyl-containing group, such as an anhydride or an acid halide, or with an alkyl group containing a good leaving group (*e.g.*, a halide) on the other.

Alternatively, it may be desirable to couple a therapeutic agent and an antibody via a linker group. A linker group can function as a spacer to distance an 10 antibody from an agent in order to avoid interference with binding capabilities. A linker group can also serve to increase the chemical reactivity of a substituent on an agent or an antibody, and thus increase the coupling efficiency. An increase in chemical reactivity may also facilitate the use of agents, or functional groups on agents, which otherwise would not be possible.

15 It will be evident to those skilled in the art that a variety of bifunctional or polyfunctional reagents, both homo- and hetero-functional (such as those described in the catalog of the Pierce Chemical Co., Rockford, IL), may be employed as the linker group. Coupling may be effected, for example, through amino groups, carboxyl groups, sulfhydryl groups or oxidized carbohydrate residues. There are numerous references 20 describing such methodology, *e.g.*, U.S. Patent No. 4,671,958, to Rodwell et al.

Where a therapeutic agent is more potent when free from the antibody portion of the immunoconjugates of the present invention, it may be desirable to use a linker group which is cleavable during or upon internalization into a cell. A number of different cleavable linker groups have been described. The mechanisms for the 25 intracellular release of an agent from these linker groups include cleavage by reduction of a disulfide bond (*e.g.*, U.S. Patent No. 4,489,710, to Spitzer), by irradiation of a photolabile bond (*e.g.*, U.S. Patent No. 4,625,014, to Senter et al.), by hydrolysis of derivatized amino acid side chains (*e.g.*, U.S. Patent No. 4,638,045, to Kohn et al.), by serum complement-mediated hydrolysis (*e.g.*, U.S. Patent No. 4,671,958, to Rodwell 30 et al.), and acid-catalyzed hydrolysis (*e.g.*, U.S. Patent No. 4,569,789, to Blattler et al.).

It may be desirable to couple more than one agent to an antibody. In one embodiment, multiple molecules of an agent are coupled to one antibody molecule. In another embodiment, more than one type of agent may be coupled to one antibody. Regardless of the particular embodiment, immunoconjugates with more than one agent 5 may be prepared in a variety of ways. For example, more than one agent may be coupled directly to an antibody molecule, or linkers that provide multiple sites for attachment can be used. Alternatively, a carrier can be used.

A carrier may bear the agents in a variety of ways, including covalent bonding either directly or via a linker group. Suitable carriers include proteins such as 10 albumins (e.g., U.S. Patent No. 4,507,234, to Kato et al.), peptides and polysaccharides such as aminodextran (e.g., U.S. Patent No. 4,699,784, to Shih et al.). A carrier may also bear an agent by noncovalent bonding or by encapsulation, such as within a liposome vesicle (e.g., U.S. Patent Nos. 4,429,008 and 4,873,088). Carriers specific for 15 radionuclide agents include radiohalogenated small molecules and chelating compounds. For example, U.S. Patent No. 4,735,792 discloses representative radiohalogenated small molecules and their synthesis. A radionuclide chelate may be formed from chelating compounds that include those containing nitrogen and sulfur atoms as the donor atoms for binding the metal, or metal oxide, radionuclide. For example, U.S. Patent No. 4,673,562, to Davison et al. discloses representative chelating 20 compounds and their synthesis.

A variety of routes of administration for the antibodies and immunoconjugates may be used. Typically, administration will be intravenous, intramuscular, subcutaneous or in the bed of a resected tumor. It will be evident that the precise dose of the antibody/immunoconjugate will vary depending upon the antibody 25 used, the antigen density on the tumor, and the rate of clearance of the antibody.

T CELLS

Immunotherapeutic compositions may also, or alternatively, comprise T cells specific for a breast tumor protein. Such cells may generally be prepared *in vitro* or *ex vivo*, using standard procedures. For example, T cells may be isolated from bone 30 marrow, peripheral blood, or a fraction of bone marrow or peripheral blood of a patient,

using a commercially available cell separation system, such as the Isolex™ System, available from Nexell Therapeutics, Inc. (Irvine, CA; see also U.S. Patent No. 5,240,856; U.S. Patent No. 5,215,926; WO 89/06280; WO 91/16116 and WO 92/07243). Alternatively, T cells may be derived from related or unrelated humans, 5 non-human mammals, cell lines or cultures.

T cells may be stimulated with a breast tumor polypeptide, polynucleotide encoding a breast tumor polypeptide and/or an antigen presenting cell (APC) that expresses such a polypeptide. Such stimulation is performed under conditions and for a time sufficient to permit the generation of T cells that are specific 10 for the polypeptide. Preferably, a breast tumor polypeptide or polynucleotide is present within a delivery vehicle, such as a microsphere, to facilitate the generation of specific T cells.

T cells are considered to be specific for a breast tumor polypeptide if the T cells specifically proliferate, secrete cytokines or kill target cells coated with the 15 polypeptide or expressing a gene encoding the polypeptide. T cell specificity may be evaluated using any of a variety of standard techniques. For example, within a chromium release assay or proliferation assay, a stimulation index of more than two fold increase in lysis and/or proliferation, compared to negative controls, indicates T cell specificity. Such assays may be performed, for example, as described in Chen et 20 al., *Cancer Res.* 54:1065-1070, 1994. Alternatively, detection of the proliferation of T cells may be accomplished by a variety of known techniques. For example, T cell proliferation can be detected by measuring an increased rate of DNA synthesis (e.g., by pulse-labeling cultures of T cells with tritiated thymidine and measuring the amount of tritiated thymidine incorporated into DNA). Contact with a breast tumor polypeptide 25 (100 ng/ml - 100 µg/ml, preferably 200 ng/ml - 25 µg/ml) for 3 - 7 days should result in at least a two fold increase in proliferation of the T cells. Contact as described above for 2-3 hours should result in activation of the T cells, as measured using standard cytokine assays in which a two fold increase in the level of cytokine release (e.g., TNF or IFN- γ) is indicative of T cell activation (see Coligan et al., Current Protocols in 30 Immunology, vol. 1, Wiley Interscience (Greene 1998)). T cells that have been activated in response to a breast tumor polypeptide, polynucleotide or polypeptide-

expressing APC may be CD4⁺ and/or CD8⁺. Breast tumor protein-specific T cells may be expanded using standard techniques. Within preferred embodiments, the T cells are derived from a patient, a related donor or an unrelated donor, and are administered to the patient following stimulation and expansion.

5 For therapeutic purposes, CD4⁺ or CD8⁺ T cells that proliferate in response to a breast tumor polypeptide, polynucleotide or APC can be expanded in number either *in vitro* or *in vivo*. Proliferation of such T cells *in vitro* may be accomplished in a variety of ways. For example, the T cells can be re-exposed to a breast tumor polypeptide, or a short peptide corresponding to an immunogenic portion
10 of such a polypeptide, with or without the addition of T cell growth factors, such as interleukin-2, and/or stimulator cells that synthesize a breast tumor polypeptide. Alternatively, one or more T cells that proliferate in the presence of a breast tumor protein can be expanded in number by cloning. Methods for cloning cells are well known in the art, and include limiting dilution.

15 PHARMACEUTICAL COMPOSITIONS

In additional embodiments, the present invention concerns formulation of one or more of the polynucleotide, polypeptide, T-cell and/or antibody compositions disclosed herein in pharmaceutically-acceptable solutions for administration to a cell or an animal, either alone, or in combination with one or more other modalities of therapy.

20 It will also be understood that, if desired, the nucleic acid segment, RNA, DNA or PNA compositions that express a polypeptide as disclosed herein may be administered in combination with other agents as well, such as, *e.g.*, other proteins or polypeptides or various pharmaceutically-active agents. In fact, there is virtually no limit to other components that may also be included, given that the additional agents do
25 not cause a significant adverse effect upon contact with the target cells or host tissues. The compositions may thus be delivered along with various other agents as required in the particular instance. Such compositions may be purified from host cells or other biological sources, or alternatively may be chemically synthesized as described herein. Likewise, such compositions may further comprise substituted or derivatized RNA or
30 DNA compositions.

Formulation of pharmaceutically-acceptable excipients and carrier solutions is well-known to those of skill in the art, as is the development of suitable dosing and treatment regimens for using the particular compositions described herein in a variety of treatment regimens, including *e.g.*, oral, parenteral, intravenous, intranasal, 5 and intramuscular administration and formulation.

1. ORAL DELIVERY

In certain applications, the pharmaceutical compositions disclosed herein may be delivered *via* oral administration to an animal. As such, these compositions may be formulated with an inert diluent or with an assimilable edible carrier, or they 10 may be enclosed in hard- or soft-shell gelatin capsule, or they may be compressed into tablets, or they may be incorporated directly with the food of the diet.

The active compounds may even be incorporated with excipients and used in the form of ingestible tablets, buccal tables, troches, capsules, elixirs, suspensions, syrups, wafers, and the like (Mathiowitz *et al.*, 1997; Hwang *et al.*, 1998; 15 U. S. Patent 5,641,515; U. S. Patent 5,580,579 and U. S. Patent 5,792,451, each specifically incorporated herein by reference in its entirety). The tablets, troches, pills, capsules and the like may also contain the following: a binder, as gum tragacanth, acacia, cornstarch, or gelatin; excipients, such as dicalcium phosphate; a disintegrating agent, such as corn starch, potato starch, alginic acid and the like; a lubricant, such as 20 magnesium stearate; and a sweetening agent, such as sucrose, lactose or saccharin may be added or a flavoring agent, such as peppermint, oil of wintergreen, or cherry flavoring. When the dosage unit form is a capsule, it may contain, in addition to materials of the above type, a liquid carrier. Various other materials may be present as coatings or to otherwise modify the physical form of the dosage unit. For instance, 25 tablets, pills, or capsules may be coated with shellac, sugar, or both. A syrup of elixir may contain the active compound sucrose as a sweetening agent methyl and propylparabens as preservatives, a dye and flavoring, such as cherry or orange flavor. Of course, any material used in preparing any dosage unit form should be pharmaceutically pure and substantially non-toxic in the amounts employed. In

addition, the active compounds may be incorporated into sustained-release preparation and formulations.

Typically, these formulations may contain at least about 0.1% of the active compound or more, although the percentage of the active ingredient(s) may, of course, be varied and may conveniently be between about 1 or 2% and about 60% or 70% or more of the weight or volume of the total formulation. Naturally, the amount of active compound(s) in each therapeutically useful composition may be prepared is such a way that a suitable dosage will be obtained in any given unit dose of the compound. Factors such as solubility, bioavailability, biological half-life, route of administration, product shelf life, as well as other pharmacological considerations will be contemplated by one skilled in the art of preparing such pharmaceutical formulations, and as such, a variety of dosages and treatment regimens may be desirable.

For oral administration the compositions of the present invention may alternatively be incorporated with one or more excipients in the form of a mouthwash, dentifrice, buccal tablet, oral spray, or sublingual orally-administered formulation. For example, a mouthwash may be prepared incorporating the active ingredient in the required amount in an appropriate solvent, such as a sodium borate solution (Dobell's Solution). Alternatively, the active ingredient may be incorporated into an oral solution such as one containing sodium borate, glycerin and potassium bicarbonate, or dispersed in a dentifrice, or added in a therapeutically-effective amount to a composition that may include water, binders, abrasives, flavoring agents, foaming agents, and humectants. Alternatively the compositions may be fashioned into a tablet or solution form that may be placed under the tongue or otherwise dissolved in the mouth.

2. INJECTABLE DELIVERY

In certain circumstances it will be desirable to deliver the pharmaceutical compositions disclosed herein parenterally, intravenously, intramuscularly, or even intraperitoneally as described in U. S. Patent 5,543,158; U. S. Patent 5,641,515 and U. S. Patent 5,399,363 (each specifically incorporated herein by reference in its entirety). Solutions of the active compounds as free base or pharmacologically acceptable salts may be prepared in water suitably mixed with a surfactant, such as

hydroxypropylcellulose. Dispersions may also be prepared in glycerol, liquid polyethylene glycols, and mixtures thereof and in oils. Under ordinary conditions of storage and use, these preparations contain a preservative to prevent the growth of microorganisms.

5 The pharmaceutical forms suitable for injectable use include sterile aqueous solutions or dispersions and sterile powders for the extemporaneous preparation of sterile injectable solutions or dispersions (U. S. Patent 5,466,468, specifically incorporated herein by reference in its entirety). In all cases the form must be sterile and must be fluid to the extent that easy syringability exists. It must be stable
10 under the conditions of manufacture and storage and must be preserved against the contaminating action of microorganisms, such as bacteria and fungi. The carrier can be a solvent or dispersion medium containing, for example, water, ethanol, polyol (e.g., glycerol, propylene glycol, and liquid polyethylene glycol, and the like), suitable mixtures thereof, and/or vegetable oils. Proper fluidity may be maintained, for
15 example, by the use of a coating, such as lecithin, by the maintenance of the required particle size in the case of dispersion and by the use of surfactants. The prevention of the action of microorganisms can be facilitated by various antibacterial and antifungal agents, for example, parabens, chlorobutanol, phenol, sorbic acid, thimerosal, and the like. In many cases, it will be preferable to include isotonic agents, for example, sugars
20 or sodium chloride. Prolonged absorption of the injectable compositions can be brought about by the use in the compositions of agents delaying absorption, for example, aluminum monostearate and gelatin.

For parenteral administration in an aqueous solution, for example, the solution should be suitably buffered if necessary and the liquid diluent first rendered
25 isotonic with sufficient saline or glucose. These particular aqueous solutions are especially suitable for intravenous, intramuscular, subcutaneous and intraperitoneal administration. In this connection, a sterile aqueous medium that can be employed will be known to those of skill in the art in light of the present disclosure. For example, one dosage may be dissolved in 1 ml of isotonic NaCl solution and either added to 1000 ml
30 of hypodermoclysis fluid or injected at the proposed site of infusion, (see for example, "Remington's Pharmaceutical Sciences" 15th Edition, pages 1035-1038 and 1570-

1580). Some variation in dosage will necessarily occur depending on the condition of the subject being treated. The person responsible for administration will, in any event, determine the appropriate dose for the individual subject. Moreover, for human administration, preparations should meet sterility, pyrogenicity, and the general safety 5 and purity standards as required by FDA Office of Biologics standards.

Sterile injectable solutions are prepared by incorporating the active compounds in the required amount in the appropriate solvent with various of the other ingredients enumerated above, as required, followed by filtered sterilization. Generally, dispersions are prepared by incorporating the various sterilized active ingredients into a 10 sterile vehicle which contains the basic dispersion medium and the required other ingredients from those enumerated above. In the case of sterile powders for the preparation of sterile injectable solutions, the preferred methods of preparation are vacuum-drying and freeze-drying techniques which yield a powder of the active ingredient plus any additional desired ingredient from a previously sterile-filtered 15 solution thereof.

The compositions disclosed herein may be formulated in a neutral or salt form. Pharmaceutically-acceptable salts, include the acid addition salts (formed with the free amino groups of the protein) and which are formed with inorganic acids such as, for example, hydrochloric or phosphoric acids, or such organic acids as acetic, 20 oxalic, tartaric, mandelic, and the like. Salts formed with the free carboxyl groups can also be derived from inorganic bases such as, for example, sodium, potassium, ammonium, calcium, or ferric hydroxides, and such organic bases as isopropylamine, trimethylamine, histidine, procaine and the like. Upon formulation, solutions will be administered in a manner compatible with the dosage formulation and in such amount 25 as is therapeutically effective. The formulations are easily administered in a variety of dosage forms such as injectable solutions, drug-release capsules, and the like.

As used herein, "carrier" includes any and all solvents, dispersion media, vehicles, coatings, diluents, antibacterial and antifungal agents, isotonic and absorption delaying agents, buffers, carrier solutions, suspensions, colloids, and the like. The use 30 of such media and agents for pharmaceutical active substances is well known in the art. Except insofar as any conventional media or agent is incompatible with the active

ingredient, its use in the therapeutic compositions is contemplated. Supplementary active ingredients can also be incorporated into the compositions.

The phrase "pharmaceutically-acceptable" refers to molecular entities and compositions that do not produce an allergic or similar untoward reaction when administered to a human. The preparation of an aqueous composition that contains a protein as an active ingredient is well understood in the art. Typically, such compositions are prepared as injectables, either as liquid solutions or suspensions; solid forms suitable for solution in, or suspension in, liquid prior to injection can also be prepared. The preparation can also be emulsified.

10 3. NASAL DELIVERY

In certain embodiments, the pharmaceutical compositions may be delivered by intranasal sprays, inhalation, and/or other aerosol delivery vehicles. Methods for delivering genes, nucleic acids, and peptide compositions directly to the lungs *via* nasal aerosol sprays has been described *e.g.*, in U. S. Patent 5,756,353 and U. S. Patent 5,804,212 (each specifically incorporated herein by reference in its entirety). Likewise, the delivery of drugs using intranasal microparticle resins (Takenaga *et al.*, 1998) and lysophosphatidyl-glycerol compounds (U. S. Patent 5,725,871, specifically incorporated herein by reference in its entirety) are also well-known in the pharmaceutical arts. Likewise, transmucosal drug delivery in the form of a polytetrafluoroethylene support matrix is described in U. S. Patent 5,780,045 (specifically incorporated herein by reference in its entirety).

4. LIPOSOME-, NANOCAPSULE-, AND MICROPARTICLE-MEDIATED DELIVERY

In certain embodiments, the inventors contemplate the use of liposomes, nanocapsules, microparticles, microspheres, lipid particles, vesicles, and the like, for the introduction of the compositions of the present invention into suitable host cells. In particular, the compositions of the present invention may be formulated for delivery either encapsulated in a lipid particle, a liposome, a vesicle, a nanosphere, or a nanoparticle or the like.

Such formulations may be preferred for the introduction of pharmaceutically-acceptable formulations of the nucleic acids or constructs disclosed herein. The formation and use of liposomes is generally known to those of skill in the art (see for example, Couvreur *et al.*, 1977; Couvreur, 1988; Lasic, 1998; which 5 describes the use of liposomes and nanocapsules in the targeted antibiotic therapy for intracellular bacterial infections and diseases). Recently, liposomes were developed with improved serum stability and circulation half-times (Gabizon and Papahadjopoulos, 1988; Allen and Choun, 1987; U. S. Patent 5,741,516, specifically incorporated herein by reference in its entirety). Further, various methods of liposome 10 and liposome like preparations as potential drug carriers have been reviewed (Takakura, 1998; Chandran *et al.*, 1997; Margalit, 1995; U. S. Patent 5,567,434; U. S. Patent 5,552,157; U. S. Patent 5,565,213; U. S. Patent 5,738,868 and U. S. Patent 5,795,587, each specifically incorporated herein by reference in its entirety).

Liposomes have been used successfully with a number of cell types that 15 are normally resistant to transfection by other procedures including T cell suspensions, primary hepatocyte cultures and PC 12 cells (Renneisen *et al.*, 1990; Muller *et al.*, 1990). In addition, liposomes are free of the DNA length constraints that are typical of viral-based delivery systems. Liposomes have been used effectively to introduce genes, drugs (Heath and Martin, 1986; Heath *et al.*, 1986; Balazsorovits *et al.*, 1989; Fresta and 20 Puglisi, 1996), radiotherapeutic agents (Pikul *et al.*, 1987), enzymes (Imaizumi *et al.*, 1990a; Imaizumi *et al.*, 1990b), viruses (Faller and Baltimore, 1984), transcription factors and allosteric effectors (Nicolau and Gersonde, 1979) into a variety of cultured cell lines and animals. In addition, several successful clinical trials examining the effectiveness of liposome-mediated drug delivery have been completed (Lopez- 25 Berestein *et al.*, 1985a; 1985b; Coune, 1988; Sculier *et al.*, 1988). Furthermore, several studies suggest that the use of liposomes is not associated with autoimmune responses, toxicity or gonadal localization after systemic delivery (Mori and Fukatsu, 1992).

Liposomes are formed from phospholipids that are dispersed in an aqueous medium and spontaneously form multilamellar concentric bilayer vesicles (also 30 termed multilamellar vesicles (MLVs). MLVs generally have diameters of from 25 nm to 4 μm . Sonication of MLVs results in the formation of small unilamellar vesicles

(SUVs) with diameters in the range of 200 to 500 Å, containing an aqueous solution in the core.

Liposomes bear resemblance to cellular membranes and are contemplated for use in connection with the present invention as carriers for the peptide 5 compositions. They are widely suitable as both water- and lipid-soluble substances can be entrapped, *i.e.* in the aqueous spaces and within the bilayer itself, respectively. It is possible that the drug-bearing liposomes may even be employed for site-specific delivery of active agents by selectively modifying the liposomal formulation.

In addition to the teachings of Couvreur *et al.* (1977; 1988), the 10 following information may be utilized in generating liposomal formulations. Phospholipids can form a variety of structures other than liposomes when dispersed in water, depending on the molar ratio of lipid to water. At low ratios the liposome is the preferred structure. The physical characteristics of liposomes depend on pH, ionic strength and the presence of divalent cations. Liposomes can show low permeability to 15 ionic and polar substances, but at elevated temperatures undergo a phase transition which markedly alters their permeability. The phase transition involves a change from a closely packed, ordered structure, known as the gel state, to a loosely packed, less-ordered structure, known as the fluid state. This occurs at a characteristic phase-transition temperature and results in an increase in permeability to ions, sugars and 20 drugs.

In addition to temperature, exposure to proteins can alter the permeability of liposomes. Certain soluble proteins, such as cytochrome c, bind, deform and penetrate the bilayer, thereby causing changes in permeability. Cholesterol inhibits this penetration of proteins, apparently by packing the phospholipids more 25 tightly. It is contemplated that the most useful liposome formations for antibiotic and inhibitor delivery will contain cholesterol.

The ability to trap solutes varies between different types of liposomes. For example, MLVs are moderately efficient at trapping solutes, but SUVs are extremely inefficient. SUVs offer the advantage of homogeneity and reproducibility in 30 size distribution, however, and a compromise between size and trapping efficiency is

offered by large unilamellar vesicles (LUVs). These are prepared by ether evaporation and are three to four times more efficient at solute entrapment than MLVs.

In addition to liposome characteristics, an important determinant in entrapping compounds is the physicochemical properties of the compound itself. Polar 5 compounds are trapped in the aqueous spaces and nonpolar compounds bind to the lipid bilayer of the vesicle. Polar compounds are released through permeation or when the bilayer is broken, but nonpolar compounds remain affiliated with the bilayer unless it is disrupted by temperature or exposure to lipoproteins. Both types show maximum efflux rates at the phase transition temperature.

10 Liposomes interact with cells *via* four different mechanisms: endocytosis by phagocytic cells of the reticuloendothelial system such as macrophages and neutrophils; adsorption to the cell surface, either by nonspecific weak hydrophobic or electrostatic forces, or by specific interactions with cell-surface components; fusion with the plasma cell membrane by insertion of the lipid bilayer of the liposome into the 15 plasma membrane, with simultaneous release of liposomal contents into the cytoplasm; and by transfer of liposomal lipids to cellular or subcellular membranes, or vice versa, without any association of the liposome contents. It often is difficult to determine which mechanism is operative and more than one may operate at the same time.

The fate and disposition of intravenously injected liposomes depend on 20 their physical properties, such as size, fluidity, and surface charge. They may persist in tissues for h or days, depending on their composition, and half lives in the blood range from min to several h. Larger liposomes, such as MLVs and LUVs, are taken up rapidly by phagocytic cells of the reticuloendothelial system, but physiology of the circulatory system restrains the exit of such large species at most sites. They can exit 25 only in places where large openings or pores exist in the capillary endothelium, such as the sinusoids of the liver or spleen. Thus, these organs are the predominate site of uptake. On the other hand, SUVs show a broader tissue distribution but still are sequestered highly in the liver and spleen. In general, this *in vivo* behavior limits the potential targeting of liposomes to only those organs and tissues accessible to their large 30 size. These include the blood, liver, spleen, bone marrow, and lymphoid organs.

Targeting is generally not a limitation in terms of the present invention. However, should specific targeting be desired, methods are available for this to be accomplished. Antibodies may be used to bind to the liposome surface and to direct the antibody and its drug contents to specific antigenic receptors located on a particular 5 cell-type surface. Carbohydrate determinants (glycoprotein or glycolipid cell-surface components that play a role in cell-cell recognition, interaction and adhesion) may also be used as recognition sites as they have potential in directing liposomes to particular cell types. Mostly, it is contemplated that intravenous injection of liposomal preparations would be used, but other routes of administration are also conceivable.

10 Alternatively, the invention provides for pharmaceutically-acceptable nanocapsule formulations of the compositions of the present invention. Nanocapsules can generally entrap compounds in a stable and reproducible way (Henry-Michelland *et al.*, 1987; Quintanar-Guerrero *et al.*, 1998; Douglas *et al.*, 1987). To avoid side effects due to intracellular polymeric overloading, such ultrafine particles (sized around 15 0.1 μm) should be designed using polymers able to be degraded *in vivo*. Biodegradable polyalkyl-cyanoacrylate nanoparticles that meet these requirements are contemplated for use in the present invention. Such particles may be easily made, as described (Couvreur *et al.*, 1980; 1988; zur Muhlen *et al.*, 1998; Zambaux *et al.* 1998; Pinto-Alphandry *et al.*, 1995 and U. S. Patent 5,145,684, specifically incorporated herein by 20 reference in its entirety).

VACCINES

In certain preferred embodiments of the present invention, vaccines are provided. The vaccines will generally comprise one or more pharmaceutical compositions, such as those discussed above, in combination with an immunostimulant. 25 An immunostimulant may be any substance that enhances or potentiates an immune response (antibody and/or cell-mediated) to an exogenous antigen. Examples of immunostimulants include adjuvants, biodegradable microspheres (*e.g.*, polylactic galactide) and liposomes (into which the compound is incorporated; *see e.g.*, Fullerton, U.S. Patent No. 4,235,877). Vaccine preparation is generally described in, for example, 30 M.F. Powell and M.J. Newman, eds., "Vaccine Design (the subunit and adjuvant

approach)," Plenum Press (NY, 1995). Pharmaceutical compositions and vaccines within the scope of the present invention may also contain other compounds, which may be biologically active or inactive. For example, one or more immunogenic portions of other tumor antigens may be present, either incorporated into a fusion 5 polypeptide or as a separate compound, within the composition or vaccine.

Illustrative vaccines may contain DNA encoding one or more of the polypeptides as described above, such that the polypeptide is generated *in situ*. As noted above, the DNA may be present within any of a variety of delivery systems known to those of ordinary skill in the art, including nucleic acid expression systems, 10 bacteria and viral expression systems. Numerous gene delivery techniques are well known in the art, such as those described by Rolland, *Crit. Rev. Therap. Drug Carrier Systems* 15:143-198, 1998, and references cited therein. Appropriate nucleic acid expression systems contain the necessary DNA sequences for expression in the patient (such as a suitable promoter and terminating signal). Bacterial delivery systems involve 15 the administration of a bacterium (such as *Bacillus-Calmette-Guerrin*) that expresses an immunogenic portion of the polypeptide on its cell surface or secretes such an epitope. In a preferred embodiment, the DNA may be introduced using a viral expression system (e.g., vaccinia or other pox virus, retrovirus, or adenovirus), which may involve the use of a non-pathogenic (defective), replication competent virus. Suitable systems are 20 disclosed, for example, in Fisher-Hoch et al., *Proc. Natl. Acad. Sci. USA* 86:317-321, 1989; Flexner et al., *Ann. N.Y. Acad. Sci.* 569:86-103, 1989; Flexner et al., *Vaccine* 8:17-21, 1990; U.S. Patent Nos. 4,603,112, 4,769,330, and 5,017,487; WO 89/01973; U.S. Patent No. 4,777,127; GB 2,200,651; EP 0,345,242; WO 91/02805; Berkner, 25 *Biotechniques* 6:616-627, 1988; Rosenfeld et al., *Science* 252:431-434, 1991; Kolls et al., *Proc. Natl. Acad. Sci. USA* 91:215-219, 1994; Kass-Eisler et al., *Proc. Natl. Acad. Sci. USA* 90:11498-11502, 1993; Guzman et al., *Circulation* 88:2838-2848, 1993; and Guzman et al., *Cir. Res.* 73:1202-1207, 1993. Techniques for incorporating DNA into such expression systems are well known to those of ordinary skill in the art. The DNA may also be "naked," as described, for example, in Ulmer et al., *Science* 259:1745-1749, 30 1993 and reviewed by Cohen, *Science* 259:1691-1692, 1993. The uptake of naked DNA may be increased by coating the DNA onto biodegradable beads, which are

efficiently transported into the cells. It will be apparent that a vaccine may comprise both a polynucleotide and a polypeptide component. Such vaccines may provide for an enhanced immune response.

It will be apparent that a vaccine may contain pharmaceutically acceptable salts of the polynucleotides and polypeptides provided herein. Such salts may be prepared from pharmaceutically acceptable non-toxic bases, including organic bases (e.g., salts of primary, secondary and tertiary amines and basic amino acids) and inorganic bases (e.g., sodium, potassium, lithium, ammonium, calcium and magnesium salts).

While any suitable carrier known to those of ordinary skill in the art may be employed in the vaccine compositions of this invention, the type of carrier will vary depending on the mode of administration. Compositions of the present invention may be formulated for any appropriate manner of administration, including for example, topical, oral, nasal, intravenous, intracranial, intraperitoneal, subcutaneous or intramuscular administration. For parenteral administration, such as subcutaneous injection, the carrier preferably comprises water, saline, alcohol, a fat, a wax or a buffer. For oral administration, any of the above carriers or a solid carrier, such as mannitol, lactose, starch, magnesium stearate, sodium saccharine, talcum, cellulose, glucose, sucrose, and magnesium carbonate, may be employed. Biodegradable microspheres (e.g., polylactate polyglycolate) may also be employed as carriers for the pharmaceutical compositions of this invention. Suitable biodegradable microspheres are disclosed, for example, in U.S. Patent Nos. 4,897,268; 5,075,109; 5,928,647; 5,811,128; 5,820,883; 5,853,763; 5,814,344 and 5,942,252. One may also employ a carrier comprising the particulate-protein complexes described in U.S. Patent No. 5,928,647, which are capable of inducing a class I-restricted cytotoxic T lymphocyte responses in a host.

Such compositions may also comprise buffers (e.g., neutral buffered saline or phosphate buffered saline), carbohydrates (e.g., glucose, mannose, sucrose or dextrans), mannitol, proteins, polypeptides or amino acids such as glycine, antioxidants, bacteriostats, chelating agents such as EDTA or glutathione, adjuvants (e.g., aluminum hydroxide), solutes that render the formulation isotonic, hypotonic or weakly hypertonic

with the blood of a recipient, suspending agents, thickening agents and/or preservatives. Alternatively, compositions of the present invention may be formulated as a lyophilizate. Compounds may also be encapsulated within liposomes using well known technology.

5 Any of a variety of immunostimulants may be employed in the vaccines of this invention. For example, an adjuvant may be included. Most adjuvants contain a substance designed to protect the antigen from rapid catabolism, such as aluminum hydroxide or mineral oil, and a stimulator of immune responses, such as lipid A, *Bordetella pertussis* or *Mycobacterium tuberculosis* derived proteins. Suitable
10 adjuvants are commercially available as, for example, Freund's Incomplete Adjuvant and Complete Adjuvant (Difco Laboratories, Detroit, MI); Merck Adjuvant 65 (Merck and Company, Inc., Rahway, NJ); AS-2 (SmithKline Beecham, Philadelphia, PA); aluminum salts such as aluminum hydroxide gel (alum) or aluminum phosphate; salts of calcium, iron or zinc; an insoluble suspension of acylated tyrosine; acylated sugars;
15 cationically or anionically derivatized polysaccharides; polyphosphazenes; biodegradable microspheres; monophosphoryl lipid A and quil A. Cytokines, such as GM-CSF or interleukin-2, -7, or -12, may also be used as adjuvants.

Within the vaccines provided herein, the adjuvant composition is preferably designed to induce an immune response predominantly of the Th1 type.
20 High levels of Th1-type cytokines (e.g., IFN- γ , TNF α , IL-2 and IL-12) tend to favor the induction of cell mediated immune responses to an administered antigen. In contrast, high levels of Th2-type cytokines (e.g., IL-4, IL-5, IL-6 and IL-10) tend to favor the induction of humoral immune responses. Following application of a vaccine as provided herein, a patient will support an immune response that includes Th1- and Th2-
25 type responses. Within a preferred embodiment, in which a response is predominantly Th1-type, the level of Th1-type cytokines will increase to a greater extent than the level of Th2-type cytokines. The levels of these cytokines may be readily assessed using standard assays. For a review of the families of cytokines, see Mosmann and Coffman, *Ann. Rev. Immunol.* 7:145-173, 1989.

30 Preferred adjuvants for use in eliciting a predominantly Th1-type response include, for example, a combination of monophosphoryl lipid A, preferably 3-

de-O-acylated monophosphoryl lipid A (3D-MPL), together with an aluminum salt. MPL adjuvants are available from Corixa Corporation (Seattle, WA; *see* US Patent Nos. 4,436,727; 4,877,611; 4,866,034 and 4,912,094). CpG-containing oligonucleotides (in which the CpG dinucleotide is unmethylated) also induce a predominantly Th1 response. Such oligonucleotides are well known and are described, for example, in WO 96/02555, WO 99/33488 and U.S. Patent Nos. 6,008,200 and 5,856,462. Immunostimulatory DNA sequences are also described, for example, by Sato et al., *Science* 273:352, 1996. Another preferred adjuvant is a saponin, preferably QS21 (Aquila Biopharmaceuticals Inc., Framingham, MA), which may be used alone or in combination with other adjuvants. For example, an enhanced system involves the combination of a monophosphoryl lipid A and saponin derivative, such as the combination of QS21 and 3D-MPL as described in WO 94/00153, or a less reactogenic composition where the QS21 is quenched with cholesterol, as described in WO 96/33739. Other preferred formulations comprise an oil-in-water emulsion and tocopherol. A particularly potent adjuvant formulation involving QS21, 3D-MPL and tocopherol in an oil-in-water emulsion is described in WO 95/17210.

Other preferred adjuvants include Montanide ISA 720 (Seppic, France), SAF (Chiron, California, United States), ISCOMS (CSL), MF-59 (Chiron), the SBAS series of adjuvants (*e.g.*, SBAS-2 or SBAS-4, available from SmithKline Beecham, Rixensart, Belgium), Detox (Corixa, Hamilton, MT), RC-529 (Corixa, Hamilton, MT) and other aminoalkyl glucosaminide 4-phosphates (AGPs), such as those described in pending U.S. Patent Application Serial Nos. 08/853,826 and 09/074,720, the disclosures of which are incorporated herein by reference in their entireties.

Any vaccine provided herein may be prepared using well known methods that result in a combination of antigen, immune response enhancer and a suitable carrier or excipient. The compositions described herein may be administered as part of a sustained release formulation (*i.e.*, a formulation such as a capsule, sponge or gel (composed of polysaccharides, for example) that effects a slow release of compound following administration). Such formulations may generally be prepared using well known technology (*see, e.g.*, Coombes et al., *Vaccine* 14:1429-1438, 1996) and administered by, for example, oral, rectal or subcutaneous implantation, or by

implantation at the desired target site. Sustained-release formulations may contain a polypeptide, polynucleotide or antibody dispersed in a carrier matrix and/or contained within a reservoir surrounded by a rate controlling membrane.

Carriers for use within such formulations are biocompatible, and may
5 also be biodegradable; preferably the formulation provides a relatively constant level of active component release. Such carriers include microparticles of poly(lactide-co-glycolide), polyacrylate, latex, starch, cellulose, dextran and the like. Other delayed-release carriers include supramolecular biovectors, which comprise a non-liquid hydrophilic core (*e.g.*, a cross-linked polysaccharide or oligosaccharide) and, optionally,
10 an external layer comprising an amphiphilic compound, such as a phospholipid (*see e.g.*, U.S. Patent No. 5,151,254 and PCT applications WO 94/20078, WO/94/23701 and WO 96/06638). The amount of active compound contained within a sustained release formulation depends upon the site of implantation, the rate and expected duration of release and the nature of the condition to be treated or prevented.

15 Any of a variety of delivery vehicles may be employed within pharmaceutical compositions and vaccines to facilitate production of an antigen-specific immune response that targets tumor cells. Delivery vehicles include antigen presenting cells (APCs), such as dendritic cells, macrophages, B cells, monocytes and other cells that may be engineered to be efficient APCs. Such cells may, but need not, be
20 genetically modified to increase the capacity for presenting the antigen, to improve activation and/or maintenance of the T cell response, to have anti-tumor effects *per se* and/or to be immunologically compatible with the receiver (*i.e.*, matched HLA haplotype). APCs may generally be isolated from any of a variety of biological fluids and organs, including tumor and peritumoral tissues, and may be autologous,
25 allogeneic, syngeneic or xenogeneic cells.

Certain preferred embodiments of the present invention use dendritic cells or progenitors thereof as antigen-presenting cells. Dendritic cells are highly potent APCs (Banchereau and Steinman, *Nature* 392:245-251, 1998) and have been shown to be effective as a physiological adjuvant for eliciting prophylactic or therapeutic
30 antitumor immunity (*see* Timmerman and Levy, *Ann. Rev. Med.* 50:507-529, 1999). In general, dendritic cells may be identified based on their typical shape (stellate *in situ*,

with marked cytoplasmic processes (dendrites) visible *in vitro*), their ability to take up, process and present antigens with high efficiency and their ability to activate naïve T cell responses. Dendritic cells may, of course, be engineered to express specific cell-surface receptors or ligands that are not commonly found on dendritic cells *in vivo* or *ex vivo*, and such modified dendritic cells are contemplated by the present invention. As an alternative to dendritic cells, secreted vesicles antigen-loaded dendritic cells (called exosomes) may be used within a vaccine (*see* Zitvogel et al., *Nature Med.* 4:594-600; 1998).

Dendritic cells and progenitors may be obtained from peripheral blood, bone marrow, tumor-infiltrating cells, peritumoral tissues-infiltrating cells, lymph nodes, spleen, skin, umbilical cord blood or any other suitable tissue or fluid. For example, dendritic cells may be differentiated *ex vivo* by adding a combination of cytokines such as GM-CSF, IL-4, IL-13 and/or TNF α to cultures of monocytes harvested from peripheral blood. Alternatively, CD34 positive cells harvested from peripheral blood, umbilical cord blood or bone marrow may be differentiated into dendritic cells by adding to the culture medium combinations of GM-CSF, IL-3, TNF α , CD40 ligand, LPS, flt3 ligand and/or other compound(s) that induce differentiation, maturation and proliferation of dendritic cells.

Dendritic cells are conveniently categorized as "immature" and "mature" cells, which allows a simple way to discriminate between two well characterized phenotypes. However, this nomenclature should not be construed to exclude all possible intermediate stages of differentiation. Immature dendritic cells are characterized as APC with a high capacity for antigen uptake and processing, which correlates with the high expression of Fc γ receptor and mannose receptor. The mature phenotype is typically characterized by a lower expression of these markers, but a high expression of cell surface molecules responsible for T cell activation such as class I and class II MHC, adhesion molecules (*e.g.*, CD54 and CD11) and costimulatory molecules (*e.g.*, CD40, CD80, CD86 and 4-1BB).

APCs may generally be transfected with a polynucleotide encoding a breast tumor protein (or portion or other variant thereof) such that the breast tumor polypeptide, or an immunogenic portion thereof, is expressed on the cell surface. Such

transfection may take place *ex vivo*, and a composition or vaccine comprising such transfected cells may then be used for therapeutic purposes, as described herein. Alternatively, a gene delivery vehicle that targets a dendritic or other antigen presenting cell may be administered to a patient, resulting in transfection that occurs *in vivo*. In 5 *vivo* and *ex vivo* transfection of dendritic cells, for example, may generally be performed using any methods known in the art, such as those described in WO 97/24447, or the gene gun approach described by Mahvi et al., *Immunology and cell Biology* 75:456-460, 1997. Antigen loading of dendritic cells may be achieved by incubating dendritic cells or progenitor cells with the breast tumor polypeptide, DNA 10 (naked or within a plasmid vector) or RNA; or with antigen-expressing recombinant bacterium or viruses (e.g., vaccinia, fowlpox, adenovirus or lentivirus vectors). Prior to loading, the polypeptide may be covalently conjugated to an immunological partner that provides T cell help (e.g., a carrier molecule). Alternatively, a dendritic cell may be pulsed with a non-conjugated immunological partner, separately or in the presence of 15 the polypeptide.

Vaccines and pharmaceutical compositions may be presented in unit-dose or multi-dose containers, such as sealed ampoules or vials. Such containers are preferably hermetically sealed to preserve sterility of the formulation until use. In general, formulations may be stored as suspensions, solutions or emulsions in oily or 20 aqueous vehicles. Alternatively, a vaccine or pharmaceutical composition may be stored in a freeze-dried condition requiring only the addition of a sterile liquid carrier immediately prior to use.

CANCER THERAPY

In further aspects of the present invention, the compositions described 25 herein may be used for immunotherapy of cancer, such as breast cancer. Within such methods, pharmaceutical compositions and vaccines are typically administered to a patient. As used herein, a "patient" refers to any warm-blooded animal, preferably a human. A patient may or may not be afflicted with cancer. Accordingly, the above pharmaceutical compositions and vaccines may be used to prevent the development of a 30 cancer or to treat a patient afflicted with a cancer. A cancer may be diagnosed using

criteria generally accepted in the art, including the presence of a malignant tumor. Pharmaceutical compositions and vaccines may be administered either prior to or following surgical removal of primary tumors and/or treatment such as administration of radiotherapy or conventional chemotherapeutic drugs. Administration may be by any 5 suitable method, including administration by intravenous, intraperitoneal, intramuscular, subcutaneous, intranasal, intradermal, anal, vaginal, topical and oral routes.

Within certain embodiments, immunotherapy may be active immunotherapy, in which treatment relies on the *in vivo* stimulation of the endogenous 10 host immune system to react against tumors with the administration of immune response-modifying agents (such as polypeptides and polynucleotides as provided herein).

Within other embodiments, immunotherapy may be passive immunotherapy, in which treatment involves the delivery of agents with established 15 tumor-immune reactivity (such as effector cells or antibodies) that can directly or indirectly mediate antitumor effects and does not necessarily depend on an intact host immune system. Examples of effector cells include T cells as discussed above, T lymphocytes (such as CD8⁺ cytotoxic T lymphocytes and CD4⁺ T-helper tumor-infiltrating lymphocytes), killer cells (such as Natural Killer cells and lymphokine-activated killer cells), B cells and antigen-presenting cells (such as dendritic cells and 20 macrophages) expressing a polypeptide provided herein. T cell receptors and antibody receptors specific for the polypeptides recited herein may be cloned, expressed and transferred into other vectors or effector cells for adoptive immunotherapy. The polypeptides provided herein may also be used to generate antibodies or anti-idiotypic 25 antibodies (as described above and in U.S. Patent No. 4,918,164) for passive immunotherapy.

Effector cells may generally be obtained in sufficient quantities for adoptive immunotherapy by growth *in vitro*, as described herein. Culture conditions for expanding single antigen-specific effector cells to several billion in number with 30 retention of antigen recognition *in vivo* are well known in the art. Such *in vitro* culture conditions typically use intermittent stimulation with antigen, often in the presence of

cytokines (such as IL-2) and non-dividing feeder cells. As noted above, immunoreactive polypeptides as provided herein may be used to rapidly expand antigen-specific T cell cultures in order to generate a sufficient number of cells for immunotherapy. In particular, antigen-presenting cells, such as dendritic, macrophage, 5 monocyte, fibroblast and/or B cells, may be pulsed with immunoreactive polypeptides or transfected with one or more polynucleotides using standard techniques well known in the art. For example, antigen-presenting cells can be transfected with a polynucleotide having a promoter appropriate for increasing expression in a recombinant virus or other expression system. Cultured effector cells for use in therapy 10 must be able to grow and distribute widely, and to survive long term *in vivo*. Studies have shown that cultured effector cells can be induced to grow *in vivo* and to survive long term in substantial numbers by repeated stimulation with antigen supplemented with IL-2 (see, for example, Cheever et al., *Immunological Reviews* 157:177, 1997).

Alternatively, a vector expressing a polypeptide recited herein may be 15 introduced into antigen presenting cells taken from a patient and clonally propagated *ex vivo* for transplant back into the same patient. Transfected cells may be reintroduced into the patient using any means known in the art, preferably in sterile form by intravenous, intracavitory, intraperitoneal or intratumor administration.

Routes and frequency of administration of the therapeutic compositions 20 described herein, as well as dosage, will vary from individual to individual, and may be readily established using standard techniques. In general, the pharmaceutical compositions and vaccines may be administered by injection (*e.g.*, intracutaneous, intramuscular, intravenous or subcutaneous), intranasally (*e.g.*, by aspiration) or orally. Preferably, between 1 and 10 doses may be administered over a 52 week period. 25 Preferably, 6 doses are administered, at intervals of 1 month, and booster vaccinations may be given periodically thereafter. Alternate protocols may be appropriate for individual patients. A suitable dose is an amount of a compound that, when administered as described above, is capable of promoting an anti-tumor immune response, and is at least 10-50% above the basal (*i.e.*, untreated) level. Such response 30 can be monitored by measuring the anti-tumor antibodies in a patient or by vaccine-dependent generation of cytolytic effector cells capable of killing the patient's tumor

cells *in vitro*. Such vaccines should also be capable of causing an immune response that leads to an improved clinical outcome (e.g., more frequent remissions, complete or partial or longer disease-free survival) in vaccinated patients as compared to non-vaccinated patients. In general, for pharmaceutical compositions and vaccines 5 comprising one or more polypeptides, the amount of each polypeptide present in a dose ranges from about 25 µg to 5 mg per kg of host. Suitable dose sizes will vary with the size of the patient, but will typically range from about 0.1 mL to about 5 mL.

In general, an appropriate dosage and treatment regimen provides the active compound(s) in an amount sufficient to provide therapeutic and/or prophylactic 10 benefit. Such a response can be monitored by establishing an improved clinical outcome (e.g., more frequent remissions, complete or partial, or longer disease-free survival) in treated patients as compared to non-treated patients. Increases in preexisting immune responses to a breast tumor protein generally correlate with an improved clinical outcome. Such immune responses may generally be evaluated using 15 standard proliferation, cytotoxicity or cytokine assays, which may be performed using samples obtained from a patient before and after treatment.

CANCER DETECTION AND DIAGNOSIS

In general, a cancer may be detected in a patient based on the presence of one or more breast tumor proteins and/or polynucleotides encoding such proteins in a 20 biological sample (for example, blood, sera, sputum urine and/or tumor biopsies) obtained from the patient. In other words, such proteins may be used as markers to indicate the presence or absence of a cancer such as breast cancer. In addition, such proteins may be useful for the detection of other cancers. The binding agents provided herein generally permit detection of the level of antigen that binds to the agent in the 25 biological sample. Polynucleotide primers and probes may be used to detect the level of mRNA encoding a tumor protein, which is also indicative of the presence or absence of a cancer. In general, a breast tumor sequence should be present at a level that is at least three fold higher in tumor tissue than in normal tissue

There are a variety of assay formats known to those of ordinary skill in 30 the art for using a binding agent to detect polypeptide markers in a sample. See, e.g.,

Harlow and Lane, *Antibodies: A Laboratory Manual*, Cold Spring Harbor Laboratory, 1988. In general, the presence or absence of a cancer in a patient may be determined by (a) contacting a biological sample obtained from a patient with a binding agent; (b) detecting in the sample a level of polypeptide that binds to the binding agent; and (c) 5 comparing the level of polypeptide with a predetermined cut-off value.

In a preferred embodiment, the assay involves the use of binding agent immobilized on a solid support to bind to and remove the polypeptide from the remainder of the sample. The bound polypeptide may then be detected using a detection reagent that contains a reporter group and specifically binds to the binding 10 agent/polypeptide complex. Such detection reagents may comprise, for example, a binding agent that specifically binds to the polypeptide or an antibody or other agent that specifically binds to the binding agent, such as an anti-immunoglobulin, protein G, protein A or a lectin. Alternatively, a competitive assay may be utilized, in which a polypeptide is labeled with a reporter group and allowed to bind to the immobilized 15 binding agent after incubation of the binding agent with the sample. The extent to which components of the sample inhibit the binding of the labeled polypeptide to the binding agent is indicative of the reactivity of the sample with the immobilized binding agent. Suitable polypeptides for use within such assays include full length breast tumor proteins and portions thereof to which the binding agent binds, as described above.

20 The solid support may be any material known to those of ordinary skill in the art to which the tumor protein may be attached. For example, the solid support may be a test well in a microtiter plate or a nitrocellulose or other suitable membrane. Alternatively, the support may be a bead or disc, such as glass, fiberglass, latex or a plastic material such as polystyrene or polyvinylchloride. The support may also be a 25 magnetic particle or a fiber optic sensor, such as those disclosed, for example, in U.S. Patent No. 5,359,681. The binding agent may be immobilized on the solid support using a variety of techniques known to those of skill in the art, which are amply described in the patent and scientific literature. In the context of the present invention, the term "immobilization" refers to both noncovalent association, such as adsorption, 30 and covalent attachment (which may be a direct linkage between the agent and functional groups on the support or may be a linkage by way of a cross-linking agent).

Immobilization by adsorption to a well in a microtiter plate or to a membrane is preferred. In such cases, adsorption may be achieved by contacting the binding agent, in a suitable buffer, with the solid support for a suitable amount of time. The contact time varies with temperature, but is typically between about 1 hour and about 1 day. In 5 general, contacting a well of a plastic microtiter plate (such as polystyrene or polyvinylchloride) with an amount of binding agent ranging from about 10 ng to about 10 µg, and preferably about 100 ng to about 1 µg, is sufficient to immobilize an adequate amount of binding agent.

Covalent attachment of binding agent to a solid support may generally be 10 achieved by first reacting the support with a bifunctional reagent that will react with both the support and a functional group, such as a hydroxyl or amino group, on the binding agent. For example, the binding agent may be covalently attached to supports having an appropriate polymer coating using benzoquinone or by condensation of an aldehyde group on the support with an amine and an active hydrogen on the binding 15 partner (see, e.g., Pierce Immunotechnology Catalog and Handbook, 1991, at A12-A13).

In certain embodiments, the assay is a two-antibody sandwich assay. This assay may be performed by first contacting an antibody that has been immobilized 20 on a solid support, commonly the well of a microtiter plate, with the sample, such that polypeptides within the sample are allowed to bind to the immobilized antibody. Unbound sample is then removed from the immobilized polypeptide-antibody complexes and a detection reagent (preferably a second antibody capable of binding to a different site on the polypeptide) containing a reporter group is added. The amount of 25 detection reagent that remains bound to the solid support is then determined using a method appropriate for the specific reporter group.

More specifically, once the antibody is immobilized on the support as described above, the remaining protein binding sites on the support are typically blocked. Any suitable blocking agent known to those of ordinary skill in the art, such as bovine serum albumin or Tween 20TM (Sigma Chemical Co., St. Louis, MO). The 30 immobilized antibody is then incubated with the sample, and polypeptide is allowed to bind to the antibody. The sample may be diluted with a suitable diluent, such as

phosphate-buffered saline (PBS) prior to incubation. In general, an appropriate contact time (*i.e.*, incubation time) is a period of time that is sufficient to detect the presence of polypeptide within a sample obtained from an individual with breast cancer. Preferably, the contact time is sufficient to achieve a level of binding that is at least about 95% of 5 that achieved at equilibrium between bound and unbound polypeptide. Those of ordinary skill in the art will recognize that the time necessary to achieve equilibrium may be readily determined by assaying the level of binding that occurs over a period of time. At room temperature, an incubation time of about 30 minutes is generally sufficient.

10 Unbound sample may then be removed by washing the solid support with an appropriate buffer, such as PBS containing 0.1% Tween 20TM. The second antibody, which contains a reporter group, may then be added to the solid support. Preferred reporter groups include those groups recited above.

15 The detection reagent is then incubated with the immobilized antibody-polypeptide complex for an amount of time sufficient to detect the bound polypeptide. An appropriate amount of time may generally be determined by assaying the level of binding that occurs over a period of time. Unbound detection reagent is then removed and bound detection reagent is detected using the reporter group. The method employed for detecting the reporter group depends upon the nature of the reporter 20 group. For radioactive groups, scintillation counting or autoradiographic methods are generally appropriate. Spectroscopic methods may be used to detect dyes, luminescent groups and fluorescent groups. Biotin may be detected using avidin, coupled to a different reporter group (commonly a radioactive or fluorescent group or an enzyme). Enzyme reporter groups may generally be detected by the addition of substrate 25 (generally for a specific period of time), followed by spectroscopic or other analysis of the reaction products.

30 To determine the presence or absence of a cancer, such as breast cancer, the signal detected from the reporter group that remains bound to the solid support is generally compared to a signal that corresponds to a predetermined cut-off value. In one preferred embodiment, the cut-off value for the detection of a cancer is the average mean signal obtained when the immobilized antibody is incubated with samples from

patients without the cancer. In general, a sample generating a signal that is three standard deviations above the predetermined cut-off value is considered positive for the cancer. In an alternate preferred embodiment, the cut-off value is determined using a Receiver Operator Curve, according to the method of Sackett et al., *Clinical Epidemiology: A Basic Science for Clinical Medicine*, Little Brown and Co., 1985, p. 106-7. Briefly, in this embodiment, the cut-off value may be determined from a plot of pairs of true positive rates (*i.e.*, sensitivity) and false positive rates (100%-specificity) that correspond to each possible cut-off value for the diagnostic test result. The cut-off value on the plot that is the closest to the upper left-hand corner (*i.e.*, the value that encloses the largest area) is the most accurate cut-off value, and a sample generating a signal that is higher than the cut-off value determined by this method may be considered positive. Alternatively, the cut-off value may be shifted to the left along the plot, to minimize the false positive rate, or to the right, to minimize the false negative rate. In general, a sample generating a signal that is higher than the cut-off value determined by this method is considered positive for a cancer.

In a related embodiment, the assay is performed in a flow-through or strip test format, wherein the binding agent is immobilized on a membrane, such as nitrocellulose. In the flow-through test, polypeptides within the sample bind to the immobilized binding agent as the sample passes through the membrane. A second, labeled binding agent then binds to the binding agent-polypeptide complex as a solution containing the second binding agent flows through the membrane. The detection of bound second binding agent may then be performed as described above. In the strip test format, one end of the membrane to which binding agent is bound is immersed in a solution containing the sample. The sample migrates along the membrane through a region containing second binding agent and to the area of immobilized binding agent. Concentration of second binding agent at the area of immobilized antibody indicates the presence of a cancer. Typically, the concentration of second binding agent at that site generates a pattern, such as a line, that can be read visually. The absence of such a pattern indicates a negative result. In general, the amount of binding agent immobilized on the membrane is selected to generate a visually discernible pattern when the biological sample contains a level of polypeptide that would be sufficient to generate a

positive signal in the two-antibody sandwich assay, in the format discussed above. Preferred binding agents for use in such assays are antibodies and antigen-binding fragments thereof. Preferably, the amount of antibody immobilized on the membrane ranges from about 25 ng to about 1 μ g, and more preferably from about 50 ng to about 5 500 ng. Such tests can typically be performed with a very small amount of biological sample.

Of course, numerous other assay protocols exist that are suitable for use with the tumor proteins or binding agents of the present invention. The above descriptions are intended to be exemplary only. For example, it will be apparent to 10 those of ordinary skill in the art that the above protocols may be readily modified to use breast tumor polypeptides to detect antibodies that bind to such polypeptides in a biological sample. The detection of such breast tumor protein specific antibodies may correlate with the presence of a cancer.

A cancer may also, or alternatively, be detected based on the presence of 15 T cells that specifically react with a breast tumor protein in a biological sample. Within certain methods, a biological sample comprising CD4 $^{+}$ and/or CD8 $^{+}$ T cells isolated from a patient is incubated with a breast tumor polypeptide, a polynucleotide encoding such a polypeptide and/or an APC that expresses at least an immunogenic portion of such a polypeptide, and the presence or absence of specific activation of the T cells is 20 detected. Suitable biological samples include, but are not limited to, isolated T cells. For example, T cells may be isolated from a patient by routine techniques (such as by Ficoll/Hypaque density gradient centrifugation of peripheral blood lymphocytes). T cells may be incubated *in vitro* for 2-9 days (typically 4 days) at 37°C with polypeptide (e.g., 5 - 25 μ g/ml). It may be desirable to incubate another aliquot of a T cell sample in 25 the absence of breast tumor polypeptide to serve as a control. For CD4 $^{+}$ T cells, activation is preferably detected by evaluating proliferation of the T cells. For CD8 $^{+}$ T cells, activation is preferably detected by evaluating cytolytic activity. A level of proliferation that is at least two fold greater and/or a level of cytolytic activity that is at least 20% greater than in disease-free patients indicates the presence of a cancer in the 30 patient.

As noted above, a cancer may also, or alternatively, be detected based on the level of mRNA encoding a breast tumor protein in a biological sample. For example, at least two oligonucleotide primers may be employed in a polymerase chain reaction (PCR) based assay to amplify a portion of a breast tumor cDNA derived from a 5 biological sample, wherein at least one of the oligonucleotide primers is specific for (*i.e.*, hybridizes to) a polynucleotide encoding the breast tumor protein. The amplified cDNA is then separated and detected using techniques well known in the art, such as gel electrophoresis. Similarly, oligonucleotide probes that specifically hybridize to a polynucleotide encoding a breast tumor protein may be used in a hybridization assay to 10 detect the presence of polynucleotide encoding the tumor protein in a biological sample.

To permit hybridization under assay conditions, oligonucleotide primers and probes should comprise an oligonucleotide sequence that has at least about 60%, preferably at least about 75% and more preferably at least about 90%, identity to a portion of a polynucleotide encoding a breast tumor protein that is at least 10 15 nucleotides, and preferably at least 20 nucleotides, in length. Preferably, oligonucleotide primers and/or probes hybridize to a polynucleotide encoding a polypeptide described herein under moderately stringent conditions, as defined above. Oligonucleotide primers and/or probes which may be usefully employed in the diagnostic methods described herein preferably are at least 10-40 nucleotides in length. 20 In a preferred embodiment, the oligonucleotide primers comprise at least 10 contiguous nucleotides, more preferably at least 15 contiguous nucleotides, of a DNA molecule having a sequence recited in SEQ ID NO: 1-38, 42-204, 205, 207 and 210-290. Techniques for both PCR based assays and hybridization assays are well known in the art (*see, for example, Mullis et al., Cold Spring Harbor Symp. Quant. Biol., 51:263, 25 1987; Erlich ed., PCR Technology, Stockton Press, NY, 1989.*)

One preferred assay employs RT-PCR, in which PCR is applied in conjunction with reverse transcription. Typically, RNA is extracted from a biological sample, such as biopsy tissue, and is reverse transcribed to produce cDNA molecules. PCR amplification using at least one specific primer generates a cDNA molecule, which 30 may be separated and visualized using, for example, gel electrophoresis. Amplification may be performed on biological samples taken from a test patient and from an

individual who is not afflicted with a cancer. The amplification reaction may be performed on several dilutions of cDNA spanning two orders of magnitude. A two-fold or greater increase in expression in several dilutions of the test patient sample as compared to the same dilutions of the non-cancerous sample is typically considered
5 positive.

In another embodiment, the compositions described herein may be used as markers for the progression of cancer. In this embodiment, assays as described above for the diagnosis of a cancer may be performed over time, and the change in the level of reactive polypeptide(s) or polynucleotide(s) evaluated. For example, the assays
10 may be performed every 24-72 hours for a period of 6 months to 1 year, and thereafter performed as needed. In general, a cancer is progressing in those patients in whom the level of polypeptide or polynucleotide detected increases over time. In contrast, the cancer is not progressing when the level of reactive polypeptide or polynucleotide either remains constant or decreases with time.

15 Certain *in vivo* diagnostic assays may be performed directly on a tumor. One such assay involves contacting tumor cells with a binding agent. The bound binding agent may then be detected directly or indirectly via a reporter group. Such binding agents may also be used in histological applications. Alternatively, polynucleotide probes may be used within such applications.

20 As noted above, to improve sensitivity, multiple breast tumor protein markers may be assayed within a given sample. It will be apparent that binding agents specific for different proteins provided herein may be combined within a single assay. Further, multiple primers or probes may be used concurrently. The selection of tumor protein markers may be based on routine experiments to determine combinations that
25 results in optimal sensitivity. In addition, or alternatively, assays for tumor proteins provided herein may be combined with assays for other known tumor antigens.

DIAGNOSTIC KITS

The present invention further provides kits for use within any of the above diagnostic methods. Such kits typically comprise two or more components
30 necessary for performing a diagnostic assay. Components may be compounds,

reagents, containers and/or equipment. For example, one container within a kit may contain a monoclonal antibody or fragment thereof that specifically binds to a breast tumor protein. Such antibodies or fragments may be provided attached to a support material, as described above. One or more additional containers may enclose elements, 5 such as reagents or buffers, to be used in the assay. Such kits may also, or alternatively, contain a detection reagent as described above that contains a reporter group suitable for direct or indirect detection of antibody binding.

Alternatively, a kit may be designed to detect the level of mRNA encoding a breast tumor protein in a biological sample. Such kits generally comprise at 10 least one oligonucleotide probe or primer, as described above, that hybridizes to a polynucleotide encoding a breast tumor protein. Such an oligonucleotide may be used, for example, within a PCR or hybridization assay. Additional components that may be present within such kits include a second oligonucleotide and/or a diagnostic reagent or container to facilitate the detection of a polynucleotide encoding a breast tumor protein.

15 The following Examples are offered by way of illustration and not by way of limitation.

EXAMPLE 1

IDENTIFICATION OF BREAST TUMOR PROTEIN cDNAS USING
SUBTRACTION METHODOLOGY

This Example illustrates the identification of cDNA molecules encoding
5 breast tumor proteins.

A human metastatic breast tumor cDNA expression library was constructed from metastatic breast tumor poly A⁺ RNA using a Superscript Plasmid System for cDNA Synthesis and Plasmid Cloning kit (BRL Life Technologies, Gaithersburg, MD 20897) following the manufacturer's protocol. Specifically, breast
10 tumor tissues were homogenized with polytron (Kinematica, Switzerland) and total RNA was extracted using Trizol reagent (BRL Life Technologies) as directed by the manufacturer. The poly A⁺ RNA was then purified using a Qiagen oligotex spin column mRNA purification kit (Qiagen, Santa Clarita, CA 91355) according to the manufacturer's protocol. First-strand cDNA was synthesized using the NotI/Oligo-
15 dT18 primer. Double-stranded cDNA was synthesized, ligated with EcoRI/BstX I adaptors (Invitrogen, Carlsbad, CA) and digested with NotI. Following size fractionation with Chroma Spin-1000 columns (Clontech, Palo Alto, CA 94303), the cDNA was ligated into the EcoRI/NotI site of pCDNA3.1 (Invitrogen, Carlsbad, CA) and transformed into ElectroMax *E. coli* DH10B cells (BRL Life Technologies) by
20 electroporation.

Using the same procedure, a normal human breast cDNA expression library was prepared from a pool of four normal breast tissue specimens. The cDNA libraries were characterized by determining the number of independent colonies, the percentage of clones that carried insert, the average insert size and by sequence analysis.
25 Sequencing analysis showed both libraries to contain good complex cDNA clones that were synthesized from mRNA, with minimal rRNA and mitochondrial DNA contamination sequencing.

A cDNA subtracted library (referred to as BS3) was prepared using the above metastatic breast tumor and normal breast cDNA libraries, as described by Hara
30 *et al.* (*Blood*, 84:189-199, 1994) with some modifications. Specifically, a breast tumor-

specific subtracted cDNA library was generated as follows. Normal breast cDNA library (70 µg) was digested with EcoRI, NotI, and SfI, followed by a filling-in reaction with DNA polymerase Klenow fragment. After phenol-chloroform extraction and ethanol precipitation, the DNA was dissolved in 100 µl of H₂O, heat-denatured and 5 mixed with 100 µl (100 µg) of Photoprobe biotin (Vector Laboratories, Burlingame, CA), the resulting mixture was irradiated with a 270 W sunlamp on ice for 20 minutes. Additional Photoprobe biotin (50 µl) was added and the biotinylation reaction was repeated. After extraction with butanol five times, the DNA was ethanol-precipitated and dissolved in 23 µl H₂O to form the driver DNA.

10 To form the tracer DNA, 10 µg breast tumor cDNA library was digested with BamHI and Xhol, phenol chloroform extracted and passed through Chroma spin-400 columns (Clontech). Following ethanol precipitation, the tracer DNA was dissolved in 5 µl H₂O. Tracer DNA was mixed with 15 µl driver DNA and 20 µl of 2 x hybridization buffer (1.5 M NaCl/10 mM EDTA/50 mM HEPES pH 7.5/0.2% sodium 15 dodecyl sulfate), overlaid with mineral oil, and heat-denatured completely. The sample was immediately transferred into a 68 °C water bath and incubated for 20 hours (long hybridization [LH]). The reaction mixture was then subjected to a streptavidin treatment followed by phenol/chloroform extraction. This process was repeated three more times. Subtracted DNA was precipitated, dissolved in 12 µl H₂O, mixed with 8 µl 20 driver DNA and 20 µl of 2 x hybridization buffer, and subjected to a hybridization at 68 °C for 2 hours (short hybridization [SH]). After removal of biotinylated double-stranded DNA, subtracted cDNA was ligated into BamHI/Xhol site of chloramphenicol resistant pBCSK⁺ (Stratagene, La Jolla, CA 92037) and transformed into ElectroMax *E. coli* DH10B cells by electroporation to generate a breast tumor specific subtracted 25 cDNA library.

To analyze the subtracted cDNA library, plasmid DNA was prepared from independent clones, randomly picked from the subtracted breast tumor specific library and characterized by DNA sequencing with a Perkin Elmer/Applied Biosystems Division Automated Sequencer Model 373A (Foster City, CA).

30 A second cDNA subtraction library containing cDNA from breast tumor subtracted with normal breast cDNA, and known as BT, was constructed as follows.

Total RNA was extracted from primary breast tumor tissues using Trizol reagent (Gibco BRL Life Technologies, Gaithersburg, MD) as described by the manufacturer. The polyA+ RNA was purified using an oligo(dT) cellulose column according to standard protocols. First strand cDNA was synthesized using the primer supplied in a Clontech 5 PCR-Select cDNA Subtraction Kit (Clontech, Palo Alto, CA). The driver DNA consisted of cDNAs from two normal breast tissues with the tester cDNA being from three primary breast tumors. Double-stranded cDNA was synthesized for both tester and driver, and digested with a combination of endonucleases (MluI, MscI, PvuII, SalI and StuI) which recognize six base pairs DNA. This modification increased the average 10 cDNA size dramatically compared with cDNAs generated according to the protocol of Clontech. The digested tester cDNAs were ligated to two different adaptors and the subtraction was performed according to Clontech's protocol. The subtracted cDNAs were subjected to two rounds of PCR amplification, following the manufacturer's protocol. The resulting PCR products were subcloned into the TA cloning vector, 15 pCRII (Invitrogen, San Diego, CA) and transformed into ElectroMax *E. coli* DH10B cells (Gibco BRL Life, Technologies) by electroporation. DNA was isolated from independent clones and sequenced using a Perkin Elmer/Applied Biosystems Division (Foster City, CA) Automated Sequencer Model 373A.

Two additional subtracted cDNA libraries were prepared from cDNA 20 from breast tumors subtracted with a pool of cDNA from six normal tissues (liver, brain, stomach, small intestine, kidney and heart; referred to as 2BT and BC6) using the PCR-subtraction protocol of Clontech, described above. A fourth subtracted library (referred to as Bt-Met) was prepared using the protocol of Clontech from cDNA from metastatic breast tumors subtracted with cDNA from five normal tissues (brain, lung, 25 PBMC, pancreas and normal breast).

cDNA clones isolated in the breast subtractions BS3, BT, 2BT, BC6 and BT-Met, described above, were colony PCR amplified and their mRNA expression levels in breast tumor, normal breast and various other normal tissues were determined using microarray technology. Briefly, the PCR amplification products were dotted onto 30 slides in an array format, with each product occupying a unique location in the array. mRNA was extracted from the tissue sample to be tested, reverse transcribed, and

fluorescent-labeled cDNA probes were generated. The microarrays were probed with the labeled cDNA probes, the slides scanned and fluorescence intensity was measured. This intensity correlates with the hybridization intensity.

The determined cDNA sequences of 131 clones determined to be over-expressed in breast tumor tissue compared to other tissues tested by a visual analysis of the microarray data are provided in SEQ ID NO: 1-35 and 42-137. Comparison of these cDNA sequences with known sequences in the gene bank using the EMBL and GenBank databases revealed no significant homologies to the sequences provided in SEQ ID NO: 7, 10, 21, 26, 30, 63, 81 and 104. The sequences of SEQ ID NO: 2-5, 8, 9, 10, 13, 15, 16, 22, 25, 27, 28, 33, 35, 72, 73, 103, 107, 109, 118, 128, 129 134 and 136 showed some homology to previously isolated expressed sequences tags (ESTs), while the sequences of SEQ ID NO: 1, 6, 11, 12, 14, 17-20, 23, 24, 29, 31, 32, 34, 42-62, 64-71, 74-80, 82-102, 105, 106, 108, 110-117, 119-127, 130-133, 135 and 137 showed some homology to previously identified genes.

15 The determined cDNA sequences of an additional 45 clones isolated from the BT-Met library as described above and found to be over-expressed in breast tumors and metastatic breast tumors compared to other tissues tested, are provided in SEQ ID NO: 138-182. Comparison of the sequences of SEQ ID NO: 159-161, 164 and 181 revealed no significant homologies to previously identified sequences. The 20 sequences of SEQ ID NO: 138-158, 162, 163, 165-180 and 182 showed some homology to previously identified genes.

In further studies, suppression subtractive hybridization (Clontech) was preformed using a pool of cDNA from 3 unique human breast tumors as the tester and a pool of cDNA from 6 other normal human tissues (liver, brain, stomach, small intestine, 25 heart and kidney) as the driver. The isolated cDNA fragments were subcloned and characterized by DNA sequencing. The determined cDNA sequences of 22 isolated clones are provided in SEQ ID NO: 183-204. Comparison of these sequences with those in the public databases revealed no significant homologies to previously identified sequences.

30 The determined cDNA sequences of 71 additional breast-specific genes isolated during characterization of breast tumor cDNA libraries are provided in SEQ ID

NO: 210-290. Comparison of these sequences with those in the GenBank and Geneseq databases revealed no significant homologies.

EXAMPLE 2

5 IDENTIFICATION OF BREAST TUMOR PROTEIN cDNAS BY RT-PCR

GABA_A receptor clones were isolated from human breast cancer cDNA libraries by first preparing cDNA libraries from breast tumor samples from different patients as described above. PCR primers were designed based on the GABA_A receptor subunit sequences described by Hedblom and Kirkness (*Jnl. Biol. Chem.* 272:15346-10 15350, 1997) and used to amplify sequences from the breast tumor cDNA libraries by RT-PCR. The determined cDNA sequences of three GABA_A receptor clones are provided in SEQ ID NO: 36-38, with the corresponding amino acid sequences being provided in SEQ ID NO: 39-41.

The clone with the longest open reading frame (ORF; SEQ ID NO: 36) 15 showed homology to the GABA_A receptor of Hedblom and Kirkness, with four potential transmembrane regions at the C-terminal part of the protein, while the clones of SEQ ID NO: 37 and 38 retained either no transmembrane region or only the first transmembrane region. Some patients were found to have only the clones with the shorter ORFs while others had both the clones with longer and shorter ORFs.

20

EXAMPLE 3

EXPRESSION OF OVARIAN TUMOR-DERIVED ANTIGENS
IN BREAST

Isolation of the antigens O772P and O8E from ovarian tumor tissue is 25 described in US Patent Application No. 09/338,933, filed June 23, 1999. The determined cDNA sequence for O772P is provided in SEQ ID NO: 205, with the corresponding amino acid sequence being provided in SEQ ID NO: 206. The full-length cDNA sequence for O8E is provided in SEQ ID NO: 207. Two protein sequences may be translated from the full length O8E. Form "A" (SEQ ID NO: 208)

begins with a putative start methionine. A second form "B" (SEQ ID NO: 209) includes 27 additional upstream residues to the 5' end of the nucleotide sequence.

The expression levels of O772P and O8E in a variety of tumor and normal tissues, including metastatic breast tumors, were analyzed by real time PCR.

5 Both genes were found to have increased mRNA expression in 30-50% of breast tumors. For O772P, elevated expression was also observed in normal trachea, ureter, uterus and ovary. For O8E, elevated expression was also observed in normal trachea, kidney and ovary. Additional analysis employing a panel of tumor cell lines demonstrated increased expression of O8E in the breast tumor cell lines SKBR3, MDA-

10 MB-415 and BT474, and increased expression of O772P in SKBR3. Collectively, the data indicate that O772P and O8E may be useful in the diagnosis and therapy of breast cancer.

EXAMPLE 4

SYNTHESIS OF POLYPEPTIDES

Polypeptides may be synthesized on a Perkin Elmer/Applied Biosystems Division 430A peptide synthesizer using FMOC chemistry with HPTU (O-Benzotriazole-N,N,N',N'-tetramethyluronium hexafluorophosphate) activation. A Gly-Cys-Gly sequence may be attached to the amino terminus of the peptide to provide a 20 method of conjugation, binding to an immobilized surface, or labeling of the peptide. Cleavage of the peptides from the solid support may be carried out using the following cleavage mixture: trifluoroacetic acid:ethanedithiol:thioanisole:water:phenol (40:1:2:2:3). After cleaving for 2 hours, the peptides may be precipitated in cold methyl-t-butyl-ether. The peptide pellets may then be dissolved in water containing 25 0.1% trifluoroacetic acid (TFA) and lyophilized prior to purification by C18 reverse phase HPLC. A gradient of 0%-60% acetonitrile (containing 0.1% TFA) in water (containing 0.1% TFA) may be used to elute the peptides. Following lyophilization of the pure fractions, the peptides may be characterized using electrospray or other types of mass spectrometry and by amino acid analysis.

From the foregoing it will be appreciated that, although specific embodiments of the invention have been described herein for purposes of illustration, various modifications may be made without deviating from the spirit and scope of the invention. Accordingly, the invention is not limited except as by the appended claims.

CLAIMS

1. An isolated polypeptide, comprising at least an immunogenic portion of a breast tumor protein, or a variant thereof that differs in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigen-specific antisera is not substantially diminished, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence selected from the group consisting of:
 - (a) sequences recited in SEQ ID NOs: 2-5, 7, 10, 13, 15, 16, 21, 22, 25, 28, 30, 33, 35, 63, 72, 73, 81, 103, 104, 107, 109, 118, 128, 129, 134, 136, 159-161, 164, 181, 183-204 and 210-290;
 - (b) sequences that hybridize to a sequence recited in any one of SEQ ID NOs: 2-5, 7, 10, 13, 15, 16, 21, 22, 25, 28, 30, 33, 35, 63, 72, 73, 81, 103, 104, 107, 109, 118, 128, 129, 134, 136, 159-161, 164, 181, 183-204 and 210-290 under moderately stringent conditions; and
 - (c) complements of sequences of (a) or (b).
2. An isolated polypeptide according to claim 1, wherein the polypeptide comprises an amino acid sequence that is encoded by a polynucleotide sequence recited in any one of SEQ ID NOs: 2-5, 7, 10, 13, 15, 16, 21, 22, 25, 28, 30, 33, 35, 63, 72, 73, 81, 103, 104, 107, 109, 118, 128, 129, 134, 136, 159-161, 164, 181, 183-204 and 210-290 or a complement of any of the foregoing polynucleotide sequences.
3. An isolated polynucleotide encoding at least 15 amino acid residues of a breast tumor protein, or a variant thereof that differs in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigen-specific antisera is not substantially diminished, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide comprising a sequence recited in any one of SEQ ID NOs: 2-5, 7, 10, 13, 15, 16, 21, 22,

25, 28, 30, 33, 35, 63, 72, 73, 81, 103, 104, 107, 109, 118, 128, 129, 134, 136, 159-161, 164, 181, 183-204 and 210-290 or a complement of any of the foregoing sequences.

4. An isolated polynucleotide encoding a breast tumor protein, or a variant thereof, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide comprising a sequence recited in any one of SEQ ID NOs: 2-5, 7, 10, 13, 15, 16, 21, 22, 25, 28, 30, 33, 35, 63, 72, 73, 81, 103, 104, 107, 109, 118, 128, 129, 134, 136, 159-161, 164, 181, 183-204 and 210-290 or a complement of any of the foregoing sequences.

5. An isolated polynucleotide, comprising a sequence recited in any one of SEQ ID NOs: 2-5, 7, 10, 13, 15, 16, 21, 22, 25, 28, 30, 33, 35, 63, 72, 73, 81, 103, 104, 107, 109, 118, 128, 129, 134, 136, 159-161, 164, 181, 183-204 and 210-290.

6. An isolated polynucleotide, comprising a sequence that hybridizes to a sequence recited in any one of SEQ ID NOs: 2-5, 7, 10, 13, 15, 16, 21, 22, 25, 28, 30, 33, 35, 63, 72, 73, 81, 103, 104, 107, 109, 118, 128, 129, 134, 136, 159-161, 164, 181, 183-204 and 210-290 under moderately stringent conditions.

7. An isolated polynucleotide complementary to a polynucleotide according to any one of claims 3-6.

8. An expression vector, comprising a polynucleotide according to any one of claims 3-7.

9. A host cell transformed or transfected with an expression vector according to claim 8.

10. An isolated antibody, or antigen-binding fragment thereof, that specifically binds to a breast tumor protein that comprises an amino acid sequence that is encoded by a polynucleotide sequence recited in any one of SEQ ID NOs: 2-5, 7, 10, 13, 15, 16, 21, 22, 25, 28, 30, 33, 35, 63, 72, 73, 81, 103, 104, 107, 109, 118, 128, 129,

134, 136, 159-161, 164, 181, 183-204 and 210-290 or a complement of any of the foregoing polynucleotide sequences.

11. A fusion protein, comprising at least one polypeptide according to claim 1.

12. A fusion protein according to claim 11, wherein the fusion protein comprises an expression enhancer that increases expression of the fusion protein in a host cell transfected with a polynucleotide encoding the fusion protein.

13. A fusion protein according to claim 11, wherein the fusion protein comprises a T helper epitope that is not present within the polypeptide of claim 1.

14. A fusion protein according to claim 11, wherein the fusion protein comprises an affinity tag.

15. An isolated polynucleotide encoding a fusion protein according to claim 11.

16. A pharmaceutical composition, comprising a physiologically acceptable carrier and at least one component selected from the group consisting of:

- (a) a polypeptide according to claim 1;
- (b) a polynucleotide according to claim 3;
- (c) an antibody according to claim 10;
- (d) a fusion protein according to claim 11; and
- (e) a polynucleotide according to claim 15.

17. A vaccine comprising an immunostimulant and at least one component selected from the group consisting of:

- (a) a polypeptide according to claim 1;

- (b) a polynucleotide according to claim 3;
- (c) an antibody according to claim 10;
- (d) a fusion protein according to claim 11; and
- (e) a polynucleotide according to claim 15.

18. A vaccine according to claim 17, wherein the immunostimulant is an adjuvant.

19. A vaccine according to any claim 17, wherein the immunostimulant induces a predominantly Type I response.

20. A method for inhibiting the development of a cancer in a patient, comprising administering to a patient an effective amount of a pharmaceutical composition according to claim 16.

21. A method for inhibiting the development of a cancer in a patient, comprising administering to a patient an effective amount of a vaccine according to claim 17.

22. A pharmaceutical composition comprising an antigen-presenting cell that expresses a polypeptide according to claim 1, in combination with a pharmaceutically acceptable carrier or excipient.

23. A pharmaceutical composition according to claim 22, wherein the antigen presenting cell is a dendritic cell or a macrophage.

24. A vaccine comprising an antigen-presenting cell that expresses a polypeptide comprising at least an immunogenic portion of a breast tumor protein, or a variant thereof, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence selected from the group consisting of:

- (a) sequences recited in SEQ ID NOs: 1-38, 42-204, 205, 207 and 210-290;
- (b) sequences that hybridize to a sequence recited in any one of SEQ ID NOs: 1-38, 42-204, 205, 207 and 210-290 under moderately stringent conditions; and
- (c) complements of sequences of (i) or (ii); in combination with an immunostimulant.

25. A vaccine according to claim 24, wherein the immunostimulant is an adjuvant.

26. A vaccine according to claim 24, wherein the immunostimulant induces a predominantly Type I response.

27. A vaccine according to claim 24, wherein the antigen-presenting cell is a dendritic cell.

28. A method for inhibiting the development of a cancer in a patient, comprising administering to a patient an effective amount of an antigen-presenting cell that expresses a polypeptide comprising at least an immunogenic portion of a breast tumor protein, or a variant thereof, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence selected from the group consisting of:

- (a) sequences recited in SEQ ID NOs: 1-38, 42-204, 205, 207 and 210-290;
- (b) sequences that hybridize to a sequence recited in any one of SEQ ID NOs: 1-38, 42-204, 205, 207 and 210-290 under moderately stringent conditions; and
- (c) complements of sequences of (i) or (ii) encoded by a polynucleotide recited in any one of SEQ ID NOs: 1-38, 42-204, 205, 207 and 210-290; and thereby inhibiting the development of a cancer in the patient.

29. A method according to claim 28, wherein the antigen-presenting cell is a dendritic cell.

30. A method according to any one of claims 20, 21 and 28, wherein the cancer is breast cancer.

31. A method for removing tumor cells from a biological sample, comprising contacting a biological sample with T cells that specifically react with a breast tumor protein, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence selected from the group consisting of:

- (i) polynucleotides recited in any one of SEQ ID NOS: 1-38, 42-204, 205, 207 and 210-290; and
- (ii) complements of the foregoing polynucleotides;

wherein the step of contacting is performed under conditions and for a time sufficient to permit the removal of cells expressing the antigen from the sample.

32. A method according to claim 31, wherein the biological sample is blood or a fraction thereof.

33. A method for inhibiting the development of a cancer in a patient, comprising administering to a patient a biological sample treated according to the method of claim 31.

34. A method for stimulating and/or expanding T cells specific for a breast tumor protein, comprising contacting T cells with at least one component selected from the group consisting of:

- (a) polypeptides comprising at least an immunogenic portion of a breast tumor protein, or a variant thereof, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence selected from the group consisting of:

(i) sequences recited in SEQ ID NOs: 1-38, 42-204, 205, 207 and 210-290;

(ii) sequences that hybridize to a sequence recited in any one of SEQ ID NOs: 1-38, 42-204, 205, 207 and 210-290 under moderately stringent conditions; and

(iii) complements of sequences of (i) or (ii);

(b) polynucleotides encoding a polypeptide of (a); and

(c) antigen presenting cells that express a polypeptide of (a); under conditions and for a time sufficient to permit the stimulation and/or expansion of T cells.

35. An isolated T cell population, comprising T cells prepared according to the method of claim 34.

36. A method for inhibiting the development of a cancer in a patient, comprising administering to a patient an effective amount of a T cell population according to claim 35.

37. A method for inhibiting the development of a cancer in a patient, comprising the steps of:

(a) incubating CD4⁺ and/or CD8⁺ T cells isolated from a patient with at least one component selected from the group consisting of:

(i) polypeptides comprising at least an immunogenic portion of a breast tumor protein, or a variant thereof, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence selected from the group consisting of:

(1) sequences recited in SEQ ID NOs: 1-38, 42-204, 205, 207 and 210-290;

(2) sequences that hybridize to a sequence recited in any one of SEQ ID NOs: 1-38, 42-204, 205, 207 and 210-290 under moderately stringent conditions; and

- (3) complements of sequences of (1) or (2);
- (ii) polynucleotides encoding a polypeptide of (i); and
- (iii) antigen presenting cells that expresses a polypeptide of (i);
 - such that T cells proliferate; and

(b) administering to the patient an effective amount of the proliferated T cells, and thereby inhibiting the development of a cancer in the patient.

38. A method for inhibiting the development of a cancer in a patient, comprising the steps of:

(a) incubating CD4⁺ and/or CD8⁺ T cells isolated from a patient with at least one component selected from the group consisting of:

- (i) polypeptides comprising at least an immunogenic portion of a breast tumor protein, or a variant thereof, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence selected from the group consisting of:
 - (1) sequences recited in SEQ ID NOs: 1-38, 42-204, 205, 207 and 210-290;
 - (2) sequences that hybridize to a sequence recited in any one of SEQ ID NOs: 1-38, 42-204, 205, 207 and 210-290 under moderately stringent conditions; and
 - (3) complements of sequences of (1) or (2);
- (ii) polynucleotides encoding a polypeptide of (i); and
- (iii) antigen presenting cells that express a polypeptide of (i);
 - such that T cells proliferate;

(b) cloning at least one proliferated cell to provide cloned T cells; and

(c) administering to the patient an effective amount of the cloned T cells, and thereby inhibiting the development of a cancer in the patient.

39. A method for determining the presence or absence of a cancer in a patient, comprising the steps of:

- (a) contacting a biological sample obtained from a patient with a binding agent that binds to a breast tumor protein, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence recited in any one of SEQ ID NOs: 1-38, 42-204, 205, 207 and 210-290 or a complement of any of the foregoing polynucleotide sequences;
- (b) detecting in the sample an amount of polypeptide that binds to the binding agent; and
- (c) comparing the amount of polypeptide to a predetermined cut-off value, and therefrom determining the presence or absence of a cancer in the patient.

40. A method according to claim 39, wherein the binding agent is an antibody.

41. A method according to claim 40, wherein the antibody is a monoclonal antibody.

42. A method according to claim 40, wherein the cancer is breast cancer.

43. A method for monitoring the progression of a cancer in a patient, comprising the steps of:

- (a) contacting a biological sample obtained from a patient at a first point in time with a binding agent that binds to a breast tumor protein, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence recited in any one of SEQ ID NOs: 1-38, 42-204, 205, 207 and 210-290 or a complement of any of the foregoing polynucleotide sequences;
- (b) detecting in the sample an amount of polypeptide that binds to the binding agent;

- (c) repeating steps (a) and (b) using a biological sample obtained from the patient at a subsequent point in time; and
- (d) comparing the amount of polypeptide detected in step (c) to the amount detected in step (b) and therefrom monitoring the progression of the cancer in the patient.

44. A method according to claim 43, wherein the binding agent is an antibody.

45. A method according to claim 44, wherein the antibody is a monoclonal antibody.

46. A method according to claim 43, wherein the cancer is a breast cancer.

47. A method for determining the presence or absence of a cancer in a patient, comprising the steps of:

- (a) contacting a biological sample obtained from a patient with an oligonucleotide that hybridizes to a polynucleotide that encodes a breast tumor protein, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence recited in any one of SEQ ID NO: 1-38, 42-204, 205, 207 and 210-290 or a complement of any of the foregoing polynucleotide sequences;

- (b) detecting in the sample an amount of a polynucleotide that hybridizes to the oligonucleotide; and

- (c) comparing the amount of polynucleotide that hybridizes to the oligonucleotide to a predetermined cut-off value, and therefrom determining the presence or absence of a cancer in the patient.

48. A method according to claim 47, wherein the amount of polynucleotide that hybridizes to the oligonucleotide is determined using a polymerase chain reaction.

49. A method according to claim 47, wherein the amount of polynucleotide that hybridizes to the oligonucleotide is determined using a hybridization assay.

50. A method for monitoring the progression of a cancer in a patient, comprising the steps of:

(a) contacting a biological sample obtained from a patient with an oligonucleotide that hybridizes to a polynucleotide that encodes a breast tumor protein, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence recited in any one of SEQ ID NO: 1-38, 42-204, 205, 207 and 210-290 or a complement of any of the foregoing polynucleotide sequences;

(b) detecting in the sample an amount of a polynucleotide that hybridizes to the oligonucleotide;

(c) repeating steps (a) and (b) using a biological sample obtained from the patient at a subsequent point in time; and

(d) comparing the amount of polynucleotide detected in step (c) to the amount detected in step (b) and therefrom monitoring the progression of the cancer in the patient.

51. A method according to claim 50, wherein the amount of polynucleotide that hybridizes to the oligonucleotide is determined using a polymerase chain reaction.

52. A method according to claim 50, wherein the amount of polynucleotide that hybridizes to the oligonucleotide is determined using a hybridization assay.

53. A diagnostic kit, comprising:

- (a) one or more antibodies according to claim 10; and
- (b) a detection reagent comprising a reporter group.

54. A kit according to claim 53, wherein the antibodies are immobilized on a solid support.

55. A kit according to claim 53, wherein the detection reagent comprises an anti-immunoglobulin, protein G, protein A or lectin.

56. A kit according to claim 53, wherein the reporter group is selected from the group consisting of radioisotopes, fluorescent groups, luminescent groups, enzymes, biotin and dye particles.

57. An oligonucleotide comprising 10 to 40 contiguous nucleotides that hybridize under moderately stringent conditions to a polynucleotide that encodes a breast tumor protein, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence recited in any one of SEQ ID NOS: 2-5, 7, 10, 13, 15, 16, 21, 22, 25, 28, 30, 33, 35, 63, 72, 73, 81, 103, 104, 107, 109, 118, 128, 129, 134, 136, 159-161, 164, 181, 183-204 and 210-290 or a complement of any of the foregoing polynucleotides.

58. An oligonucleotide according to claim 57, wherein the oligonucleotide comprises 10-40 contiguous nucleotides recited in any one of SEQ ID NOS: 2-5, 7, 10, 13, 15, 16, 21, 22, 25, 28, 30, 33, 35, 63, 72, 73, 81, 103, 104, 107, 109, 118, 128, 129, 134, 136, 159-161, 164, 181, 183-204 and 210-290.

59. A diagnostic kit, comprising:
(a) an oligonucleotide according to claim 58; and
(b) a diagnostic reagent for use in a polymerase chain reaction or hybridization assay.

SEQUENCE LISTING

<110> Corixa Corporation
Dillon, Davin C.
Day, Craig H.
Jiang, Yuqiu
Wang, Aijun
Houghton, Raymond L.
Mitcham, Jennifer L.

<120> COMPOSITIONS AND METHODS FOR THERAPY AND
DIAGNOSIS OF BREAST CANCER

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agacacagca taaaactgaa agcaccatgg cataaaagtct agtaacatca tcctcaaaag      420
cttttccaa tgccttcct tcaactgttt attcagtatt tggccagtag aaataaagat      480
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ctgcatggtt tcttatattgc aagcacaaga catggcacat tggttccact gtacaggttag	180
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gcttcttana aaataganaa cttcaatgg tcataataca tttgattca aaatgtcttc	300
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<210> 15	
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<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
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gaaagcaaaa ctgcaaaaca tagtcttgg cattcacatt tgcttcagca gtataattaa	300
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<211> 547	
<212> DNA	
<213> Homo sapien	
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<223> n = A,T,C or G	
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gagaatactg ccaggcttt cctaattctt ttggctttt gaaatggca gggttctca	180
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aagacacttt tagccaatga agttttcaaa agaagaaagc ctctgttgcgcttttttgc	180
atatgcactg aacttctgaa atatcttttcc caaaaagtcc acaaatttctt tttccaaatc	240
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<210> 19	
<211> 239	
<212> DNA	
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<212> DNA							
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aaggatgcga	atcaggactt	ggttaattggg	ctcagttcc	tagtagggga	agaaagagat	180	
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<213> Homo sapien							
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catgactaca	tacagtacat	cctacaggca	aagagagg	gaagggggaaa	aagaagactg	180	
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<210> 28

<211> 530

<212> DNA

<213> Homo sapien

<400> 28

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<210> 29

<211> 571

<212> DNA

<213> Homo sapien

<220>

<221> misc_feature

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<400> 29

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aaatatattt attatgcact ttcataataca cagggatttt ttgagtagcca angggataaa	420
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<210> 30

<211> 917

<212> DNA

<213> Homo sapien

<400> 30

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<210> 31
 <211> 367
 <212> DNA
 <213> Homo sapien

<220>
 <221> misc_feature
 <222> (1)...(367)
 <223> n = A,T,C or G

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<210> 32
 <211> 847
 <212> DNA
 <213> Homo sapien

<400> 32	
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<210> 33
 <211> 863
 <212> DNA
 <213> Homo sapien

<220>
 <221> misc_feature
 <222> (1)...(863)
 <223> n = A,T,C or G

<400> 33

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<210> 34

<211> 432

<212> DNA

<213> Homo sapien

<400> 34

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<210> 35

<211> 350

<212> DNA

<213> Homo sapien

<400> 35

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<210> 36

<211> 1082

<212> DNA

<213> Homo sapien

<400> 36

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tggggctatg atggaaatga tgtggagttc acctggctga gagggAACGA ctctgtgcgt	600
ggactggAACAC acctgcggct tgctcagtac accatAGAGC ggtatttcac cttAGTCACC	660
agatcgcAGC aggAGACAGG aaATTACACT agATTGGTCT tacAGTTGA gttcggagg	720
aatgttctgt atttcatttt gatatctct cgattcagtc cctgcaagaa cctgcattgg	780
ggacaacaaa ggaAGTAGAA gaAGTCAGTA ttACTAATAT catcaACAGC tccatctcca	840
gcttaaacg gaAGATCAGC tttGCCAGCA ttGAAATTTC cAGCGACAAC gttgactaca	900
gtgacttgac aatgAAAACC agcgacaAGT taaAGTTGT cttccgagaa aagatgggca	960
ggattgttga ttATTCACA attcaAAACC ccAGTAATGT tgatcaCTAT tccAAACTAC	1020
tgtttcTTT gatTTTATG ctAGCCAATG tattttactg ggcataCTAC atgtatTTT	1080
ga	1082

<210> 37
<211> 1135
<212> DNA
<213> Homo sapien

<400> 37	
atgaactaca gcctccactt ggcTTcgtg tgtctgagtc tcttcactga gaggatgtgc	60
atccaggggatc gtcagttcaa cgTCGAGGTC ggcagaAGTG acaagcttcc cctgcctggc	120
tttggaaacc tcacAGCAGG atataacaaa ttTCTCAGGC ccaattttgg tgGAGAACCC	180
gtacAGATAG cgCTGACTCT ggACATTGCA agtatctcta gcatttcaga gagtaACATG	240
gactACACAG ccACCATATA CCTCCGACAG cgCTGGATGG accAGCGGCT ggtGTTGAA	300
ggcaacaAGA gcttcactct ggATGCCGc cTCGTGGAGT tccTCTGGGT gccAGATACT	360
taCATTGTGG agtccaAGAA gtcTTTCCtC catgaAGTCA ctgtGGGAAA caggCTCATC	420
cgcctttctt ccaatggcac ggtcctgtat gccctcagaa tcacgacaac tttgcatgt	480
aacatggatc tgtctaaata ccccatggac acacagacat gcaagttgca gctggaaagc	540
tggggctatg atggaaatga tgtggagttc acctggctga gagggAACGA ctctgtgcgt	600
ggactggAACAC acctgcggct tgctcagtac accatAGAGC ggtatttcac cttAGTCACC	660
agatcgcAGC aggAGACAGG aaATTACACT agATTGGTCT tacAGTTGA gttcggagg	720
aatgttctgt atttcatttt gatatctctt cttccTTGAGT ggtGTTGTCC	780
tggTTTcat tttggatCTC tCTCGATTCA gtcCCTGCAA gaACCCGcat tgGGGACAAC	840
aaAGGAAGTA gaAGAAGTCA gtattactaa tatcatcaac agtCCATCT ccAGCTTAA	900
acGGAAGATC agCTTGCcA gCATTGAAAT ttCCAGCGAC AACGTTGACT acAGTgACTT	960
gacaatgAAA accAGCGACA agttaaAGTT tgcTTCCGA gaaaAGATGG gCAGGATTGT	1020
tgattatttc acaattcaAA accCCAGTAA tgcTTGATCAC tattccAAAC tactgtttcc	1080
tttGATTTT atgctAGCCA atgtatTTA ctggcatcc tacatgtatt ttGA	1135

<210> 38
<211> 1323
<212> DNA
<213> Homo sapien

<400> 38	
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atccaggggatc gtcagttcaa cgTCGAGGTC ggcagaAGTG acaagcttcc cctgcctggc	120
tttggaaacc tcacAGCAGG atataacaaa ttTCTCAGGC ccaattttgg tgGAGAACCC	180
gtacAGATAG cgCTGACTCT ggACATTGCA agtatctcta gcatttcaga gagtaACATG	240
gactACACAG ccACCATATA CCTCCGACAG cgCTGGATGG accAGCGGCT ggtGTTGAA	300
ggcaacaAGA gcttcactct ggATGCCGc cTCGTGGAGT tccTCTGGGT gccAGATACT	360
taCATTGTGG agtccaAGAA gtcTTTCCtC catgaAGTCA ctgtGGGAAA caggCTCATC	420
cgcctttctt ccaatggcac ggtcctgtat gccctcagaa tcacgacaac tttgcatgt	480
aacatggatc tgtctaaata ccccatggac acacagacat gcaagttgca gctggaaagc	540
tggggctatg atggaaatga tgtggagttc acctggctga gagggAACGA ctctgtgcgt	600
ggactggAACAC acctgcggct tgctcagtac accatAGAGC ggtatttcac cttAGTCACC	660
agatcgcAGC aggAGACAGG aaATTACACT agATTGGTCT tacAGTTGA gttcggagg	720
aatgttctgt atttcatttt gatatctctt cttccTTGAGT ggtGTTGTCC	780

tgggttcat ttggatctc ttcgattca gtccctgcaa gaacctgcac tggagtgacg 840
accgtttat caatgaccac actgatgatc gggccccca cttctttcc caacaccaac 900
tgcttcatca agccatcgta tgttacctg gggatctgtt ttagcttgc gtttggggcc 960
ttgctagaat atgcagttgc tcactacagt tccttacagc agatggcagc caaagatagg 1020
gggacaacaa aggaagtaga agaagtcagt attactaata tcatcaacag ctccatctcc 1080
agctttaaac ggaagatcag ctttgccagc attgaaattt ccagcgacaa cgttgactac 1140
agtgacttga caatgaaaac cagcgacaag ttcaagtttgc tcttccgaga aaagatggc 1200
aggattgttgc attatttcac aattcaaaac cccagtaatg ttgatcacta ttccaaacta 1260
ctgtttccct tgatTTTtat gctagccaat gtatTTTact gggcatacta catgtatTTT 1320
tga 1323

<210> 39
<211> 440
<212> PRT
<213> *Homo sapien*

<400> 39
 Met Asn Tyr Ser Leu His Leu Ala Phe Val Cys Leu Ser Leu Phe Thr
 1 5 10 15
 Glu Arg Met Cys Ile Gln Gly Ser Gln Phe Asn Val Glu Val Gly Arg
 20 25 30
 Ser Asp Lys Leu Ser Leu Pro Gly Phe Glu Asn Leu Thr Ala Gly Tyr
 35 40 45
 Asn Lys Phe Leu Arg Pro Asn Phe Gly Gly Glu Pro Val Gln Ile Ala
 50 55 60
 Leu Thr Leu Asp Ile Ala Ser Ile Ser Ser Ile Ser Glu Ser Asn Met
 65 70 75 80
 Asp Tyr Thr Ala Thr Ile Tyr Leu Arg Gln Arg Trp Met Asp Gln Arg
 85 90 95
 Leu Val Phe Glu Gly Asn Lys Ser Phe Thr Leu Asp Ala Arg Leu Val
 100 105 110
 Glu Phe Leu Trp Val Pro Asp Thr Tyr Ile Val Glu Ser Lys Lys Ser
 115 120 125
 Phe Leu His Glu Val Thr Val Gly Asn Arg Leu Ile Arg Leu Phe Ser
 130 135 140
 Asn Gly Thr Val Leu Tyr Ala Leu Arg Ile Thr Thr Thr Val Ala Cys
 145 150 155 160
 Asn Met Asp Leu Ser Lys Tyr Pro Met Asp Thr Gln Thr Cys Lys Leu
 165 170 175
 Gln Leu Glu Ser Trp Gly Tyr Asp Gly Asn Asp Val Glu Phe Thr Trp
 180 185 190
 Leu Arg Gly Asn Asp Ser Val Arg Gly Leu Glu His Leu Arg Leu Ala
 195 200 205
 Gln Tyr Thr Ile Glu Arg Tyr Phe Thr Leu Val Thr Arg Ser Gln Gln
 210 215 220
 Glu Thr Gly Asn Tyr Thr Arg Leu Val Leu Gln Phe Glu Leu Arg Arg
 225 230 235 240
 Asn Val Leu Tyr Phe Ile Leu Glu Thr Tyr Val Pro Ser Thr Phe Leu
 245 250 255
 Val Val Leu Ser Trp Val Ser Phe Trp Ile Ser Leu Asp Ser Val Pro
 260 265 270
 Ala Arg Thr Cys Ile Gly Val Thr Thr Val Leu Ser Met Thr Thr Leu
 275 280 285
 Met Ile Gly Ser Arg Thr Ser Leu Pro Asn Thr Asn Cys Phe Ile Lys
 290 295 300
 Ala Ile Asp Val Tyr Leu Gly Ile Cys Phe Ser Phe Val Phe Gly Ala
 305 310 315 320
 Leu Leu Glu Tyr Ala Val Ala His Tyr Ser Ser Leu Gln Gln Met Ala

	325	330	335
Ala Lys Asp Arg Gly Thr Thr Lys Glu Val Glu Val Ser Ile Thr			
340	345	350	
Asn Ile Ile Asn Ser Ser Ile Ser Ser Phe Lys Arg Lys Ile Ser Phe			
355	360	365	
Ala Ser Ile Glu Ile Ser Ser Asp Asn Val Asp Tyr Ser Asp Leu Thr			
370	375	380	
Met Lys Thr Ser Asp Lys Phe Lys Phe Val Phe Arg Glu Lys Met Gly			
385	390	395	400
Arg Ile Val Asp Tyr Phe Thr Ile Gln Asn Pro Ser Asn Val Asp His			
405	410	415	
Tyr Ser Lys Leu Leu Phe Pro Leu Ile Phe Met Leu Ala Asn Val Phe			
420	425	430	
Tyr Trp Ala Tyr Tyr Met Tyr Phe			
435	440		

<210> 40
 <211> 289
 <212> PRT
 <213> Homo sapien

	<400> 40		
Met Asn Tyr Ser Leu His Leu Ala Phe Val Cys Leu Ser Leu Phe Thr			
1	5	10	15
Glu Arg Met Cys Ile Gln Gly Ser Gln Phe Asn Val Glu Val Gly Arg			
20	25	30	
Ser Asp Lys Leu Ser Leu Pro Gly Phe Glu Asn Leu Thr Ala Gly Tyr			
35	40	45	
Asn Lys Phe Leu Arg Pro Asn Phe Gly Gly Glu Pro Val Gln Ile Ala			
50	55	60	
Leu Thr Leu Asp Ile Ala Ser Ile Ser Ser Ile Ser Glu Ser Asn Met			
65	70	75	80
Asp Tyr Thr Ala Thr Ile Tyr Leu Arg Gln Arg Trp Met Asp Gln Arg			
85	90	95	
Leu Val Phe Glu Gly Asn Lys Ser Phe Thr Leu Asp Ala Arg Leu Val			
100	105	110	
Glu Phe Leu Trp Val Pro Asp Thr Tyr Ile Val Glu Ser Lys Lys Ser			
115	120	125	
Phe Leu His Glu Val Thr Val Gly Asn Arg Leu Ile Arg Leu Phe Ser			
130	135	140	
Asn Gly Thr Val Leu Tyr Ala Leu Arg Ile Thr Thr Thr Val Ala Cys			
145	150	155	160
Asn Met Asp Leu Ser Lys Tyr Pro Met Asp Thr Gln Thr Cys Lys Leu			
165	170	175	
Gln Leu Glu Ser Trp Gly Tyr Asp Gly Asn Asp Val Glu Phe Thr Trp			
180	185	190	
Leu Arg Gly Asn Asp Ser Val Arg Gly Leu Glu His Leu Arg Leu Ala			
195	200	205	
Gln Tyr Thr Ile Glu Arg Tyr Phe Thr Leu Val Thr Arg Ser Gln Gln			
210	215	220	
Glu Thr Gly Asn Tyr Thr Arg Leu Val Leu Gln Phe Glu Leu Arg Arg			
225	230	235	240
Asn Val Leu Tyr Phe Ile Leu Glu Thr Tyr Val Pro Ser Thr Phe Leu			
245	250	255	
Val Val Leu Ser Trp Val Ser Phe Trp Ile Ser Leu Asp Ser Val Pro			
260	265	270	
Ala Arg Thr Arg Ile Gly Asp Asn Lys Gly Ser Arg Arg Ser Gln Tyr			
275	280	285	

Tyr

<210> 41
<211> 265
<212> PRT
<213> Homo sapien

<400> 41

Met	Asn	Tyr	Ser	Leu	His	Leu	Ala	Phe	Val	Cys	Leu	Ser	Leu	Phe	Thr
1				5					10					15	
Glu	Arg	Met	Cys	Ile	Gln	Gly	Ser	Gln	Phe	Asn	Val	Glu	Val	Gly	Arg
				20				25					30		
Ser	Asp	Lys	Leu	Ser	Leu	Pro	Gly	Phe	Glu	Asn	Leu	Thr	Ala	Gly	Tyr
	35					40						45			
Asn	Lys	Phe	Leu	Arg	Pro	Asn	Phe	Gly	Gly	Glu	Pro	Val	Gln	Ile	Ala
	50					55				60					
Leu	Thr	Leu	Asp	Ile	Ala	Ser	Ile	Ser	Ser	Ile	Ser	Glu	Ser	Asn	Met
	65					70				75				80	
Asp	Tyr	Thr	Ala	Thr	Ile	Tyr	Leu	Arg	Gln	Arg	Trp	Met	Asp	Gln	Arg
					85			90					95		
Leu	Val	Phe	Glu	Gly	Asn	Lys	Ser	Phe	Thr	Leu	Asp	Ala	Arg	Leu	Val
		100				105			105				110		
Glu	Phe	Leu	Trp	Val	Pro	Asp	Thr	Tyr	Ile	Val	Glu	Ser	Lys	Lys	Ser
		115				120			120			125			
Phe	Leu	His	Glu	Val	Thr	Val	Gly	Asn	Arg	Leu	Ile	Arg	Leu	Phe	Ser
		130				135			135			140			
Asn	Gly	Thr	Val	Leu	Tyr	Ala	Leu	Arg	Ile	Thr	Thr	Thr	Val	Ala	Cys
	145					150				155			160		
Asn	Met	Asp	Leu	Ser	Lys	Tyr	Pro	Met	Asp	Thr	Gln	Thr	Cys	Lys	Leu
					165				170			175			
Gln	Leu	Glu	Ser	Trp	Gly	Tyr	Asp	Gly	Asn	Asp	Val	Glu	Phe	Thr	Trp
		180				185			185			190			
Leu	Arg	Gly	Asn	Asp	Ser	Val	Arg	Gly	Leu	Glu	His	Leu	Arg	Leu	Ala
		195				200			200			205			
Gln	Tyr	Thr	Ile	Glu	Arg	Tyr	Phe	Thr	Leu	Val	Thr	Arg	Ser	Gln	Gln
		210				215			215			220			
Glu	Thr	Gly	Asn	Tyr	Thr	Arg	Leu	Val	Leu	Gln	Phe	Glu	Leu	Arg	Arg
	225					230			230			235			240
Asn	Val	Leu	Tyr	Phe	Ile	Leu	Asp	Leu	Ser	Arg	Phe	Ser	Pro	Cys	Lys
					245			245			250			255	
Asn	Leu	His	Trp	Gly	Gln	Gln	Arg	Lys							
					260			265							

<210> 42
<211> 574
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(574)
<223> n = A,T,C or G

<400> 42

accaacanag	cttagtaatt	tctaaaaaga	aaaaatgatc	ttttccgac	ttctaaacaa	60
gtgactatac	tagcataaat	cattcttcta	gtaaaacagc	taaggtatag	acattctaat	120
aatttggaa	aacctatgat	tacaagtaaa	aactcagaaa	tgcaaagatg	ttggttttt	180

gtttctcagt ctgcttagc ttttaactct ggaaacgcgt gcacactgaa ctctgctcag
 tgctaaacag tcaccagcag gttcctcagg gttcagccc taaaatgtaa aacctggata
 atcagtgtat gttgcaccag aatcagcatt ttttttttaa ctgaaaaaaaa tgatggtctc
 atctctgaat ttatatttct cattctttg aacatactat agtaatata ttttatgttg
 ctaaatgtct tctatctagc atgttaaaca aagataaat actttcgatg aaagtaaatt
 ataggaaaaaa aattaactgt tttaaaaaga acttgattat gttttatgtat ttcaggcaag
 tattcatttt taacttgcta cctactttta aata

<210> 43
 <211> 467
 <212> DNA
 <213> Homo sapien

<220>
 <221> misc_feature
 <222> (1) ... (467)
 <223> n = A,T,C or G

<400> 43
 tttttttttt ttttttattt ccatcaattt attaaaataa acatgtataag caggtttcaa
 caattgtctt gtagtttcta gtaaaaaagac ataagaaaaga gaaggtgtgg tttgcagcaa
 tccgttagctg gtttctcacc ataccctgca gttctgtgag ccaaagggtct tgcagaaaagt
 taaaataaaat cacaaagact gctgtcatat attaattgca taaacacacctc aacattgctc
 anagtttcat ccgtttgggt aanaaaaacat tccttcaattt catctatggc attttagtgc
 gcattgtcgat ctatgaactc ttgaagaagt tctttgtattt cagtcttaga cacttggaa
 ttgattgtct tggaaatcac attctccaat aaggggcagc cagagcctgc gtagcagtgc
 tgggagaggg ccgccagcat gaggaccatc agcaacttca tggttag
 <210> 44
 <211> 613
 <212> DNA
 <213> Homo sapien

<220>
 <221> misc_feature
 <222> (1) ... (613)
 <223> n = A,T,C or G

<400> 44
 tttttttttt ttttttttag ttttaaaaata ttttcaacttt attattatgc ttataatatt
 attccaacag actgtattaa aggcaagtat cactaacaca gaacacgaca gggcgaagag
 gcagccgggc cgattgcagg acgtggcctg tcgggccagg gtcgctgaca tgcacgctgg
 tagctcatac actgctaccc tcagcacagg ctgcaggaat agggacaaga cagatgccgc
 cggactctta gaagcttattt aataaatatc atccaaaaac aaaatggaaa agaaacaaga
 aaccctccga gcacaaccac cttaggccaa ctgaatgtaa tctagttat tcaaccaaaa
 attgagagag aaggaaaata ttgaacacaa caaacgaaag aaagcagttc ttaagactag
 cagtaaataaa atttatacaa cagttcggtc tgtataatat gatgaaataa atctacatct
 tttcttattt tggngctttg aattatacat acaaacaaca attacaggga cttgttcaca
 aagcatgttag gcctanaaaaa aggctctctg aaaccctcaa tggcaactgg tgaacggtaa
 cactgattgc cca

<210> 45
 <211> 334
 <212> DNA
 <213> Homo sapien

<220>
 <221> misc feature

<222> (1) . . . (334)
 <223> n = A,T,C or G

 <400> 45
 accagaccaa gtgaatgcga caggaaatta tttcctgtgt tgataattca tgaagttagaa 60
 cagtataatc aaaatcaatt gtatcatcat tagtttcca ctgcctcaca ctatgtgagct 120
 gtgccaagta gtagtgtgac acctgtgttg tcatttcca catcacgtaa gagcttccaa 180
 gaaaaagccaa atcccagatg agtctcagag agggatcaat atgtccatga ttatcaggta 240
 tgctgactat ttccaagggg ttttcagtt gcttcatttg cttgtaaagc aggtaatcct 300
 cttgttgtnt tttcttttc tcgatgagcc gtgt 334

 <210> 46
 <211> 429
 <212> DNA
 <213> Homo sapien

 <220>
 <221> misc_feature
 <222> (1) . . . (429)
 <223> n = A,T,C or G

 <400> 46
 acaatttnt taaacaagca gaatagcact aggccagaata aaaaattgca cagacgtatg 60
 caattttcca agatagcatt cttaaattc agtattcagc ttccaaagat tggttgccc 120
 taatagactt aaacatataa tgatggctaa aaaaaataag tatacgaaaa tgtaaaaaag 180
 gaaatgttaag tccactctca atctcataaa aggtgagagt aaggatgcta aagcaaaata 240
 aatgttagtt cttttttctt atttccgttt atcatgcagt ctgcttctt gatatgcctt 300
 agggttaccc atttaagttt gagggtgtaa tgcaatggtg ggaatgaaaa ttgatcaa 360
 atacaccttg tcatttcatt tcaaattgcg gntggaaact tccaaaaaaaaa gggtaggcat 420
 gaagaaaaaa 429

 <210> 47
 <211> 394
 <212> DNA
 <213> Homo sapien

 <220>
 <221> misc_feature
 <222> (1) . . . (394)
 <223> n = A,T,C or G

 <400> 47
 acgcgaantt gtgttatgac tgatagcctt cagctacaaa angataggac tgacctggtt 60
 taaagtgttc tattttgtaa atcattccat ttgagtcattt ctgatgaact tggctatact 120
 gaaatctgtt atttttagtga ggctccaaaa tgagccaaagc taggcctgtat tagagtagag 180
 tgactattaa aaaacataac tttcttaggag ctataaatca aagttttaaa aagatgtttg 240
 gatatatttgcg agtattccga tcatgaaaac agaaattgcc ctgcctacta caaggacaga 300
 ctgatggaa attatgcacc tggtaactt agcttttaag cagacgatgc tgtaaaaaca 360
 aacggcttctt ctgatattta ttgtaagttt tagt 394

 <210> 48
 <211> 486
 <212> DNA
 <213> Homo sapien

 <400> 48
 acaaaggaac cgaggggtga ccacctctga gatgtcctt actttgtcat agcctggggc 60
 atattgagca tctctctcac agctgcctt cttatccccca ttcttgatgt agacccctt 120

ccgagtcagc ttttctcct cctcagacac aaacagagct ttgatatacct gtgcagggag	180
cagcttcc ttttgtgt ggcaagtgggt agttggagga agcctcaaag ctgcagttgt	240
tccctcggtg caggggagac aaatgggcct gatagtctgg ccatatttca gcttatttctt	300
gagcttgatc agggcaacgt catagtcata aaattcagga attcctgctt ctttttccc	360
attaatgttg tagttgggtt gaaataggac tacttctatc tccaggtccc gcttctcccc	420
tcccttgatt gagtgttccct tgtcatccac agtgaacaa tgcgtgcgt tcagcacaaa	480
gtacct	486

<210> 49
<211> 487
<212> DNA
<213> Homo sapien

<400> 49

acgggctgac agagaagatt cccgagagta aatcatctt ccaatccaga ggaacaagca	60
tgtctctctg ccaagatcca tctaaactgg agtgcgttgc gcaagacccag ctttagagttc	120
ttctttcttt cttaaaggccct ttgcgtctgg ggaagttctc cagttcagc tcaactcaca	180
gcttcctccaa gcatcaccct gggagttcc tgagggtttt ctcataaaatg agggctgcac	240
attgcctgtt ctgcgtcgaa gtattcaata ccgcgtca ttttaaatga agtgcgttca	300
agatttggtt tggatcaat aggaaagcat atgcagccaa ccaagatgca aatgtttga	360
aatgatatac cccaaaattt aagtaggaaa gtcacccaaa cacttctgtt ttcacttaag	420
tgtctggccc gcaatactgtt aggaacaagc atgatctgtt tactgtgata ttttaaatat	480
ccacagt	487

<210> 50
<211> 460
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(460)
<223> n = A,T,C or G

<400> 50

acatattttt gttgaagacca ccagactgaa gtaaacagct gtgcacccaa tttattatag	60
ttttgttaatg aacaatatgt aatcaaaactt ctaggtgact tgagagtggaa acctcctata	120
tcattattta gcaccgtta tgacagtaac catttcagtg tattgtttat tataccactt	180
atatacaactt attttcacc aggttaaat ttaatttct acaaaaataac attctgaatc	240
aagcacactg tatgttcagt aggttgaact atgaacactg tcatcaatgt tcagttcaaa	300
agcctgaaag tttagatcta gaagctggta aaaatgacaa tatcaatcac attagggaa	360
ccattgtgtt cttcaactaa tccatattgc actattgaaa ataaggcacac caagntatat	420
gactaatata acttgaaaat ttttataact gaggggtng	460

<210> 51
<211> 529
<212> DNA
<213> Homo sapien

<400> 51

acacttgaaa ccaaatttctt aaaacttggtt tttcttaaaa aatagttgtt gtaacattaa	60
accataacct aatcgtgtt ttcactatgc ttccacacta gccagtcttc tcacacttct	120
tctggttca agtctcaagg cctgacagac agaaggctt ggagattttt tttctttaca	180
attcagtctt cagcaacttgc agagtttctt tcatgttgc aagcaacaga gctgtatctg	240
cagggtcgta agcatagaga cggttgaat atcttcaggat gatatcggtt ctaactgtca	300
gagatgggtc aacaaacata atcctgggaa catactggcc atcaggagaa aggtgtttgt	360
cagttgttcc ataaaccaga ttgaggagga caaaactgttc tgccaaatttc tggatttctt	420
tatatttcaggc aacactttc tttaaagctt gactgtgtgg gcactcatcc aagtgtgaa	480

taaatcatca	agggtttgtt	gcttgtcttg	gatttatata	gagcttctt	529	
<210> 52						
<211> 379						
<212> DNA						
<213> Homo sapien						
<400> 52						
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acatgaacgt	tttca	tc gcttca	gcctggagat	ctgcttcaga	gaaatcttg	60
tttggact	caaaagtatg	tccagaaaat	cccagcgct	tttctgagta	gtatcttgg	120
ttagtttac	c ttaa	aggagac	tc ttccgg	cctggattac	tttctctgtg	180
gttcttggtt	aaat	tttagaa	a agat	tttgc	aactgatgaa	240
tgtgatgt	aaaattgttc	atgcgctggt	tggagat	ttt gctt	gctaagg	300
tcaggtatga	gtccagg	gtt	ttt	ttt	ttt	360
						379
<210> 53						
<211> 380						
<212> DNA						
<213> Homo sapien						
<220>						
<221> misc_feature						
<222> (1)...(380)						
<223> n = A,T,C or G						
<400> 53						
acttttatct	taaaagggtg	gtagtttcc	ctaaaatact	tattatgtaa	gggtcattag	60
acaatgtct	tgaagt	tagac	atggaattt	tgaatgg	tttattcattt	120
ttttggcat	cctgg	tttgc	ctccagttt	aggc	tttgc	180
ggaacac	ctg	gggggc	tcttcc	tttgc	tttgc	240
tttgaaatt	ataga	aaattt	actatgtaa	tgcttgatgg	aatnnttcc	300
gcttctgaaa	ggcg	ctttct	ccatttattt	aaaactaccc	atgcaattaa	360
gccgcgacca	cnctaang	gc				380
<210> 54						
<211> 245						
<212> DNA						
<213> Homo sapien						
<400> 54						
gcgcggcgct	tcacttcttc	aacttccgg	ccggctcgcc	cagcgcgt	cgagtgt	60
ccgagg	tgca	ggagg	ccgc	gcgt	ggat	120
tctt	cagcac	agagcg	cac	tttacttca	tttgc	180
ggaaatgttc	tgctcg	agg	tttcaaga	atcaga	aaacc	240
cttgt						245
<210> 55						
<211> 556						
<212> DNA						
<213> Homo sapien						
<400> 55						
acagaagatg	aataataatg	aaaaactgtg	atttttgac	tatcacatac	attgtgtt	60
aaaacaggta	aatataatga	cttactgt	taagaaagac	aaggaggaaa	actgttt	120
tgttcagg	tttactaa	gcacaaaaat	ataacaaatt	ctgtgt	tac	180
gaagtgtata	caagtgcatt	gcaaatgagc	tctt	aaaat	ttaaagtcca	240
agccaagcat	atgtctacat	ttatgattt	tttctt	ttt	ttt	300

agtttttaa aaagttcat catggctgtc atcttgaat ctgcctcca gctcaaagct	360
gagacttcac gcatacatat ttcctttctt ggttgcatt tcaccttagtt tctccaaagta	420
ttcagagttt aataggcaca cttcttttat atgttcaatt ttgtccacat gtatggcag	480
tgctgctgct tcagtaggct ttctcacaca ccctttccct tcttcaaca gcagtccacca	540
aacgttcaca acacaa	556
<210> 56	
<211> 166	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1) ... (166)	
<223> n = A,T,C or G	
<400> 56	
atggccctg attacatcat tatgaactac tcaggnnaac atccaaata ccgacctngg	60
gaaagacttg gtccgagatg ttttcatcca tacaggtac ctttccaga gcncaggnc	120
caagagctgc ntatcacct acctggccca ggtggacccc anaggg	166
<210> 57	
<211> 475	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1) ... (475)	
<223> n = A,T,C or G	
<400> 57	
acatccncat gttcctccaa atgacgttg gggcctgtc tgccaacatt ctttattgcc	60
agctgttcag gtgtcatctt atcttcttct tctacgcct tattgtatt cttggctaat	120
tccaaacatct ctttaccac tgattcattg cgtttacaat gttcactgtc gtcctgaagt	180
gtcaaacctt ccatccaaact cttttatgc aaatttagca acatcttctg ttccagttca	240
ttttccat agttaataatg aatggagtaa taatgtctgt ttagtccatg aattaatgcc	300
tggatagatg gcttgtttaa gtgacccaga ttcgaagttg tttgtcttg ttcatgtcct	360
aagaccatca tattagcatt gatcaatctg aaggcatcaa taacaaccctt tcctttaca	420
ctctgaatgg gatccacaaac cactgccaca gntctctccg ataaggcttc aaagc	475
<210> 58	
<211> 520	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1) ... (520)	
<223> n = A,T,C or G	
<400> 58	
actgttnatg tgctacttgc atttgtccct cttcctgtgc actaaagacc ccactcaatt	60
cccttagtgtt cagcagtggta tgacctctag tcaagacatt tgcacttagga tagttatgt	120
gaaccatggc aactgatcac aacaatgtct ttcagatcg atccattttt tcctcttgc	180
tttacagcaa gggatattaa ttacctatgt taccttccc tggactatg aatgtgcaaa	240
attccaatgt tcatggcttc tccctttaaa cctatattct acccccttta cattatagaa	300
aggaatgctg gaaacccaga gtccttctct tggactctt aatgtgtatt tctaattatc	360

catgactctt aatgtgcata tttcaattg cctaatngat ttcaattgtc taagacattt	420
caaagtcta attggggaga actgagtctt ttatatcaag ctaatatcta gcttttat	480
caagctaata tcttgacttc tcagcatcat agaagggggt	520
<210> 59	
<211> 214	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(214)	
<223> n = A,T,C or G	
<400> 59	
ctggcaggaa atgcatcaaa agacttaaag gtanagcgta ttaccctcg tcacttgcaa	60
cggcttattc gtggagatga agaattggat tctctcatca aggctacaat tgctgggtgn	120
ggtgtcattc cacacatcca caaatctctg atnnggaana aaggacaaca naagactgnc	180
taanggatgc ctgnatncc tgaatctca tgac	214
<210> 60	
<211> 360	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(360)	
<223> n = A,T,C or G	
<400> 60	
gcataacaaca tggcagcagg gcctcgaa ganggttagg aggaccgagc agcattctct	60
gttagaggaag acaggaaagg agaccctttt ggcacacatt tatggaggggt tgccttgaa	120
gagaaggca ggtggagag gtccctgtt acttaagaga aggcaccagt ggcaaaagagc	180
acaatgaaga ggatgtatgt aaaaacaatc acgcagataa ggacaatcat cttcacgttc	240
ttccaccaga atttcgagc caccttctgc gatgtcgctc tgaagtgctc agatgtggct	300
tccagatcct ctgtttgtt gcggagatgt tccaaatccc cccccgggc caggatccgc	360
<210> 61	
<211> 391	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(391)	
<223> n = A,T,C or G	
<400> 61	
tntggatcg tactcgatta aacagagcca ctttggttcc tgaggcaatg cataantcan	60
cattttcaa tgactgcttc ttttggaaag gnttggagat gactttatc cgcttgctga	120
ggaacacacc aatgnccatca ctgttgcattt agaacatctt tacagacaac atgaantgct	180
ttcgcttgtc tgagtcaatg atatacaatg ttttggctgt gcaatagttc tttccttcca	240
agtttagctg ctgcatttct tggncactat ttccatccc aataaatgca cacgggtttag	300
actcttgcata agaacaacca tcnccgttcca tttgttcttt ttttnttcttc catccactgc	360
ccataagata tacacannga ggtggcaaa a	391
<210> 62	

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<211> 324
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(324)
<223> n = A,T,C or G

<400> 62
acaattttat tttaacagat ttcaggagtc catttttaa aaaatgagca ataaagaacc      60
tctatcagtg agacttctca tttatagca aatacattt tgcagctaa attttcttga      120
attcatatac gcttctgtca tttaaacaaa cttccagaga aaactggct ctatataattt      180
aagtaacaaa ttgacaaaa tacatattt tacatatata ganctctaata ataaatatta      240
aatttgaaaa aatcaaatgt gaagcagaaa ctgctataca agtatattgt ntaatatcta      300
tntnatacat taaagnnttc cggg      324

<210> 63
<211> 360
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(360)
<223> n = A,T,C or G

<400> 63
acagannct tgaatatgtt gtggttccct cattatggcc cttcattccc ttctgtgtta      60
atagtaaagc atgttgctta ataactacaa ccctgaccaa atttgggcct ggatctcatg      120
ggtcacgtgg agttttaaat acgatttttta atttacttgg gtaattgagc tgaatcttta      180
gttttcagat tactttttta aacagatagg ctcttagaac aaatttattaa aaacataata      240
ccccatttggaa gggaaatctg gattaactac ccactgttcc cacccccccc aactttgaa      300
aaattttggc catatagaat gcatgaaaaaa tcaggtatga tcttatgagg actttatagt      360

<210> 64
<211> 491
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(491)
<223> n = A,T,C or G

<400> 64
nctgactgtg atgtccactt gttccctgat ttttacacat catgtcaaag ataacagctg      60
ttcccaccca ccagttctc taagcacata ctctgtttt ctgtcaacat cccattttgg      120
ggaaaggaaa agtcatattt attcccgac cccagttttt taacttggttc tcccagttgt      180
ccccctcttc tctgggtgta agaaggggaaa ttggaaaaaa attatatata tatttcctt      240
ttaatggtgg ggggctactg gagaggagag acagcaagtc caccctaact tgttacacag      300
cacataccac aggttctgga attctcatct tcgaacctag agaaataggt gctataaaca      360
ggaaattaag caaatgtg gatgtatag atcttttaat tgncttaatt tttttctat      420
tattaaacta caggctgttag atntcttagg tctcacagaa ctntatcat tttaaactga      480
cttgtatatt t      491

<210> 65
<211> 484

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<212> DNA
 <213> Homo sapien

<220>
 <221> misc_feature
 <222> (1)...(484)
 <223> n = A,T,C or G

<400> 65

accagcacac	cggcgccgtc	ctggactgct	ccttctacga	tccaacgcac	gcctggagt	60
gaggactaga	tcatcaattt	aaaatgcac	atttgaacac	tgatcaagaa	aatcttgtt	120
ggaccatga	tgccccatc	agatgtgt	aatactgtcc	agaagtgaat	gtgtatggta	180
ctggaaatgt	ggatcagaca	gctaaactgt	gggatcccag	aactccttgt	aatgctggg	240
ccttctctca	gcctgaaaag	gtatataccc	tctcagtgtc	tggagaccgg	ctgattgtgg	300
gaacagcagg	ccgcagagng	ttggtgtggg	acttacggaa	catgggttac	gtgcagcagg	360
gcagggagtc	cagcctgaaa	taccagactc	gctgcatacg	agcgtttcca	aacaaggcagg	420
gttatgtatt	aagcttattt	gaaggccgag	tggcagttga	gtatttggac	ccaagccctg	480
aggt						484

<210> 66
 <211> 355
 <212> DNA
 <213> Homo sapien

<220>
 <221> misc_feature
 <222> (1)...(355)
 <223> n = A,T,C or G

<400> 66

ngaagaaaat	atgggtggag	gtgaaggtaa	tcacagagct	gctgatttcc	aaaacagtgg	60
tgaaggaaat	acagggtctg	cagaatcttc	tttttctcag	gaggtttcta	gagaacaaca	120
gccatcatca	gcatctgaaa	gacaggcccc	tcgagcacct	cagtcaccga	gacgcccacc	180
acatccactt	cccccaagac	tgaccattca	tgccccacct	caggagttgg	gaccaccagt	240
tcagagaatt	catatgaccc	gaaggcagtc	tgttaggacgt	ggccttcagt	tgactccagg	300
aataggtggc	acgcaacacgc	attttttga	tgtatgaagac	agaacagttc	caagt	355

<210> 67
 <211> 417
 <212> DNA
 <213> Homo sapien

<400> 67

acgacacccc	tcaagagggt	gccgaagctt	tcctgtcttc	cctgacagag	accatagaag	60
gagtcgtgc	tgaggatggg	cacagcccg	gggaacaaca	gaagcggaaag	atcgctctgg	120
acccttcagg	ctccatgaac	atctacctgg	tgctagatgg	atcagacacgc	attggggcca	180
gcaacttcac	aggagccaaa	aagtgtctag	tcaacttaat	tgagaagggt	gcaagttatg	240
gtgtgaagtc	aagatatggt	ctagtgcacat	atgcacata	ccccaaaatt	tgggtcaaag	300
tgtctgaagc	agacagcagt	aatgcagact	gggtcacgaa	gcagctaat	gaaatcaatt	360
atgaagacca	caagttgaag	tcagggacta	acaccaagaa	ggccctccag	gcagtgt	417

<210> 68
 <211> 223
 <212> DNA
 <213> Homo sapien

<220>
 <221> misc_feature

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<222> (1)...(223)
<223> n = A,T,C or G

<400> 68
cactgcaag cttgcttaca gagacctgn t aaacaaagaa cagacagatt ctataaaatc      60
agtatatca acatataaag gagtgtgatt ttcagttgt tttttaagt aaatatgacc      120
aaactgacta aataagaagg caaaacaaaa aattatgctt ccttgacaag gccttggag      180
taaacaaaat gcttaaggc tcctggtaa tgggttgca agg      223

<210> 69
<211> 396
<212> DNA
<213> Homo sapien

<400> 69
acctttttc tctccaaagg aacagttctt aaagtttctt gggggaaaaaaa aaaacttaca      60
tcaaatttaa accatatgtt aaactgcata tttagttgt tacaccaaaa aattgcctca      120
gctgatctac acaagttca aagtctttaa tgcttgatataaatttactc aacattaaat      180
tatcttaat tattaattaa aaaaaaaaaact ttcttaaggaa aaataaacaa attagaccc      240
tgattatcaa aggattatta aagaatctt accaaaaatt tcaaccctac aacctaaaaac      300
cgcaaatttc tatTTTaaa catcagaaaa taactcttgg ttcattactt atgacccaaa      360
gttttattt cactattcaa tatctgaaaa gtatca      396

<210> 70
<211> 402
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(402)
<223> n = A,T,C or G

<400> 70
acccannccc acccaggcaa acagctccga catgttngt aagttagagaca agccagtgca      60
agttttttt tttttttctt ttttctttt tttgtttttt gcttaccttc ttgtttaatg      120
gaattgttat ggctaagcac atagaaggcc aaaaaaggag ttttcaaac ccagcaaattc      180
aagtgttgg attctgaact gccaaaaagaa aactgcactt cccctctttaa gtaaaacgaa      240
atgagttctt tagttaatg tattcatcag cccagataaa aaaaaaaccg gttatgtgag      300
cgtagtcac tgctcatttc caggaanatc aaacaaaata ccagcccagc cagactcaca      360
tgtgggnata tatataaaa gcaagagagc cacacccaca ag      402

<210> 71
<211> 385
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(385)
<223> n = A,T,C or G

<400> 71
accagttagag agtggccct gcaggccact tataaacagg aagctctctc ctgagctcac      60
tgatcaacct gcccggca cagacagaac ctaccagaaa agaacaagta caaaaacta      120
tcattatctg ttttctcaag acagtccaa atgtcctgt gcgatcgcca caaactcagt      180
gattggccca agtcattccc ggggccata aacagtaact ggtgtgcanc attagaacaa      240
ggggacacgg ctttgattctt cttctgagca acatgaactg ggatttctgc cnccccggat      300

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ctcggctgcc acctccgaag aagtctgtac cagccacctc cacagtaaaa gattcctccc	360
gtgagttatga tttggaaatgc gnccct	385
<210> 72	
<211> 538	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(538)	
<223> n = A,T,C or G	
<400> 72	
caattaatta acagaggat aattgtctca ctttcagaag tgatcatatata tttttatata	60
gcacaggta taagaaaaat atatagaaaa ataatcaatt tcataatataa aaggattatt	120
tctccacatt taattattgg cctatcattt gttagtgtt tttggcata ttattgaact	180
aatgtattat tccattcaaa gtctttctag attaaaaat gtatgaaaaa gcttaggatt	240
atatacatgtg taactattat agataaacatc ctaaaccttc agtttagata tataattgac	300
tgggtgtaat ctctttgtt atctgnnttg acagattttt taaattatgt tagcataatc	360
aaggaagatt taccttgaag cacttccaa attgataactt tcaaacttat tttaaagcag	420
tagaaccttt tctatgaact aagtccatcg caaaactcca acctgttaagt atacataaaa	480
tggacttact tattcctctc accttctcca ggccttagaa tattctctc tggagccc	538
<210> 73	
<211> 405	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(405)	
<223> n = A,T,C or G	
<400> 73	
actttatnna tgaaatttcc ttctacttgt atccattncc cggggcttat ggaccattc	60
atactctcca tatttagaat caaagggtcc ttctgaaga gaccttaatt ttaaggtaaa	120
acgtggtcca agttcctgaa ttcccactt ctttcactc ctgaatatgt atctgtgaaa	180
tctgaagaat atgtatccc gtgtattgtt gaatgtggca acctgccttc cgataaaattg	240
aggattatga ggaaagagag atgcaaacat acgtccattt gaatgaccca gccgtgttgt	300
aaaatttattt agaatttattt caggtatgtt ttctgtgggg tccttgcttc ttctcttaat	360
ttcttacga agacgaacac tgctcatttt aaaatgagca gttgg	405
<210> 74	
<211> 498	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(498)	
<223> n = A,T,C or G	
<400> 74	
tgagccctgc acctgtttcc tgacccccctt gccnactggc tctatggcca caaggagttt	60
taccctgtttaa aggagtttga ggtgtattat aagctgtatgg aaaaataccc atgtgtgtt	120
ccctgtggg ttggaccctt tacgtatgttcc ttctgtgtcc atgacccaga ctatgccaag	180
attctcctga aaagacaaga tccaaaatgtt gctgttagcc acaaaaatctt tgaatcctgg	240

gttggtcgag gacttgcac cctggatgg tctaaatgga aaaagcacccg ccagattgt	300
aaacctggct tcaacatcg cattctgaaa atattcatca ccatgtatgc tgagagtgtt	360
cggatgtgc tgaacaaatg ggaggaacac attgccccaa actcacgtct ggagctttt	420
caacatgtct ccctgtatgc cctggacagc atcatgaagt gtgcctttag ccaccaggc	480
agcatccagt tggacagt	498
<210> 75	
<211> 458	
<212> DNA	
<213> Homo sapien	
<400> 75	
agccttgcac atgatactca gattcctcac cttgttag gagtaaaaca atatactta	60
cagggtgata ataatctcca tagttatgg aagtggctt aaaaaggcaa gattgacttt	120
tatgacattt gataaaatct acaaattcagc cctcgagttt ttcaatgata actgacaaac	180
taaatttattt ccctagaaag gaagatgaaa ggagtggagt gtggtttggc agaacaactg	240
catttcacag ctttccagt taaattggag cactgaacgt tcagatgtt accaaattat	300
gcatgggtcc taatcacaca tataaggctg gctaccagct ttgacacagc actgttcatc	360
tggccaaaca actgtggta aaaacacatg taaaatgctt tttaacagct gataactgtat	420
aagacaaagc caagatgcaa aattaggctt tgattggc	458
<210> 76	
<211> 340	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(340)	
<223> n = A,T,C or G	
<400> 76	
accttataacc aaaanaatgc ttattccaaa atatttttg tagcttagtag ttcttcctt	60
ggaggttaaag aaaatacacc caaacttttta attaccagga ttcaagaatat ttaagagaac	120
aatttttagtt aagaatcaaa tatactgaga ttcaagagg ggaaaaaaag gaaatattat	180
agaagacaaa ggtcaaactg gcattccaga tctggagca ttttgtaaag cagaaaaaca	240
actatgacaa tctgnagctt cttagatcat tatagtaat gtncccat tataaaggg	300
tttttataat ggttttcctt aaataaagga acataaatgt	340
<210> 77	
<211> 405	
<212> DNA	
<213> Homo sapien	
<400> 77	
actccatttg tggaaactcg tgcggagtct ggttaaacagc cgaatgtctt cctccctac	60
agtttcctct cttgcataa gagcagtgtt gtcctgatta aaggcattaa ttttatctat	120
caggaagaac attttttcat ttctgttcc cggtatgtcg acaccatact tttgtagctc	180
ctctgttatt ctctggtag ttccttgcatt ttgattttctt aacaggggca gagattaca	240
gatatgtgtg atgagctcgc tggtaagttt ttctgcagg cagggaaaccg tggcccttcc	300
ttcctccagc agatccctga aatatgggtt gttctcaaag aagatcttct ctctctgcag	360
ggcttcggac aggctcagct ggtcctggat ctctgttgg ccccg	405
<210> 78	
<211> 410	
<212> DNA	
<213> Homo sapien	

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<220>
<221> misc_feature
<222> (1)...(410)
<223> n = A,T,C or G

<400> 78
acagcagntn tagatggctg caacaacattt cctccttaccc cagcccagaa aatatttctg      60
ccccacccca ggatccggga cccaaaataaa gagcaagcag gcccccttca ctgaggtgct      120
ggtagggct cagtgcacca ttactgtgct ttgagaaaga ggaaggggat ttgtttggca      180
ctttaaaaat agaggagtaa gcaggactgg agaggccaga gaagatacca aaattggcag      240
ggagagacca tttggcgcca gtccccctagg agatgggagg agggagatag gtatgagggt      300
aggcgctaag aagagttagga ggggtccact ccaagtggca gggtgctgaa atgggctagg      360
accaacagga cactgactct agttttatga cctgtccata cccgttccac      410

<210> 79
<211> 512
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(512)
<223> n = A,T,C or G

<400> 79
acagtgaaaa acaaactaat ataaaggcatt ccagnngata aaaaccttcc caggcttatg      60
gttttttc caaggaaattt atgttcaat gtaaagtttgg aaatactcca gacatacatt      120
ccatgttagt tttgggtgcc aatgttaaaa tttcaaaattt tgcatgcaag gcttagcaaa      180
gaaacactgg cagaattcca gcatttgcaa aattctaagt tttggtgaat attgttaata      240
ttacaattgg tattagaaag ccatgatgaa tccagaatta agagaaaacc catttcataa      300
atatttgtt tgattaaaaa ataccaggct taccatgttc taaataaacac aagaaaatat      360
ctttaaaaaa aaaaggactg caatttaaca gtaatctgta tatcttttagc tgcattaaa      420
aaaagaaaaa agaacaacca aaaacaatga aaatgttaca actggtataa agtnaccnna      480
tgatgctccc cttacgagaa aacaaaactg tc      512

<210> 80
<211> 174
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(174)
<223> n = A,T,C or G

<400> 80
tgattccca gacctaataat gggctaacac gcttctttc tncagcagnc ttccctgtccg      60
tgaagntncc ttccagattt gtacatggaa ctgaaaacaa agggagcctc agctggattt      120
aaatctggag catgccacaa agncttgcac tnggcatttt cnagaagaac ccat      174

<210> 81
<211> 274
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(274)

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<223> n = A,T,C or G

<400> 81

ttgcaacaag cacattaaat taaggcctgc tngaatttct tcctcccaa tcaggtaaac	60
tttccttgcc aataaagttt gaggaggtgg catttggaaa tctctttaaa aaagaagtct	120
tcatctattc acnagaaaac tcaaaaataa ttttcattat caacacacaa actaactcaa	180
tctctgcttt aagtttctat tggccaattt ttctgattna tacgagaatt attntcagnt	240
ntagaaaatc ctggtcttg gtcattacaa gntg	274

<210> 82

<211> 101

<212> DNA

<213> Homo sapien

<220>

<221> misc_feature

<222> (1)...(101)

<223> n = A,T,C or G

<400> 82

atggagaaga tcgaacctga gcctnnntgag aattgcctgc tacngcctgg cagccctgcc	60
cgagtggccc agcnncattt cacnagntgg gcatgatttg n	101

<210> 83

<211> 182

<212> DNA

<213> Homo sapien

<400> 83

tattatgggg aaagataact gagaataaag ctatcatgca gatatttgc a gataaaaag	60
taatgcagat actgagtgg a gtttgc aactatgctt gaaagccact ctaccactag	120
ttacacaaac caataatttc ctttcgcagt ggaagtcagc ttgagtttt tcaggtgttt	180
tt	182

<210> 84

<211> 229

<212> DNA

<213> Homo sapien

<220>

<221> misc_feature

<222> (1)...(229)

<223> n = A,T,C or G

<400> 84

actgtttgta gctgcactac aacagattct taccgtctcc acaaaggta gagattgtaa	60
atggtaata ctgactttt ttttattccc ttgactcaag acagctaact tcattttcag	120
aactgtttta aacctttgtg tgctggttta taaaataatg tggtaatcc ttgttgcttt	180
cctgatacca nactgttcc cgnggttggt tagaatatat tnngttcng	229

<210> 85

<211> 500

<212> DNA

<213> Homo sapien

<220>

<221> misc_feature

<222> (1)...(500)

<223> n = A,T,C or G

<400> 85

ggggagtang tgatttatta aagcaagacg ttgaaacctt tacnttctgc agtgaagatc	60
agggtgtcat tgaaagacag tggaaaccag gatgaaagtt tttacatgtc acacactaca	120
tttcttcataat atttcacca ggacttccgc aatgaggctt cgtttctgaa gggacatctg	180
atccgagcat ctcttcactc ctaacttggc tgcaacagct tccagagggg catcaaattt	240
ggcaagactt aacttgaaca gaggttcaact aatgaagaag aagtctaaca gctcagaaac	300
aagagctggg cagaactcg cattggctg gttagcagcag agggccagcg tgaccagcag	360
gagacacacc gacagctca tggggcttg ttttgcgtg agctcagctt tcacaaacaa	420
tgagtgattt ggactccacc ccaggagcct gtggagctgc agagcccagg gctatttga	480
cctgccccggg cggncgctcg	500

<210> 86

<211> 323

<212> DNA

<213> Homo sapien

<220>

<221> misc_feature

<222> (1)...(323)

<223> n = A,T,C or G

<400> 86

ccgccagtgt gctggaaattc gcccttgccg cccgggcagg tactcagaag tcatttgtt	60
tttacaattt ggttgtgtg ggatggatn tanggcggat gagccagtgc ttttgcatt	120
aagatcaat antcattgtc ctctccact gtctccctt tcctcacccc atggcagctn	180
tcatgaccca ttcccaaagg gtccaccgag tcctgaactc agcttcatca ccaacattcc	240
tcgccttcag ttgaattcaa cactgncaan ggagnagang caaagacttg ggtcagggag	300
agggngggaa acacanaaca aac	323

<210> 87

<211> 230

<212> DNA

<213> Homo sapien

<400> 87

gcagcattga gccacccctt tggcaggcga tacggcagct ctgtccctt gcccagcatg	60
tggagtggag gagatgtgc ccctgtggtt ggaacatcct ggggtgaccc cgcacccagc	120
ctcgctggc tgtccccgtt ccctatctt cactctggac ccaggctga catcctaata	180
aaataactgt tggatttagacaaaaaaaaaaaaaaa aaaaaaaaaaagg	230

<210> 88

<211> 249

<212> DNA

<213> Homo sapien

<220>

<221> misc_feature

<222> (1)...(249)

<223> n = A,T,C or G

<400> 88

atgtgaccag gtctaggctt ggagttcag ntggacact gagccaagca gacaagcaaa	60
gcaaggccagg acacaccatc ctgccccagg cccagcttctt ctcctgcctt ccaacgccc	120
ggggagcaat ctcagccccc aactctgcctt gatgcctttt atcttggggc tcttgcgtt	180
aggtgtgacc accactccnt ggtcttggc ccggcccat ggatcctgct ctctggaggg	240
ggtnatgt	249

<210> 89
 <211> 203
 <212> DNA
 <213> Homo sapien

<220>
 <221> misc_feature
 <222> (1)...(203)
 <223> n = A,T,C or G

<400> 89
 tgtttacact gtcaaggatg acaaggaaag tgttcntatc tntgatacca tcatacccagc 60
 tgttcctcct cccactgacc tgcgattcac caacattgg ccagacacca tgcgtgtcac 120
 ctgggctcca cccccatcta ttgatttaac taacttcctg gtgcgnnact cacctgtcaa 180
 aaatgangaa gatgttgcag agt 203

<210> 90
 <211> 455
 <212> DNA
 <213> Homo sapien

<400> 90
 ctctaagggg gctggcaaca tggctcagca ggcttgcccc agagccatgg caaagaatgg 60
 acttgtaatt tgcacatcctgg tgatcacctt actcctggac cagaccacca gccacacatc 120
 cagattaaaa gccaggaagc acagcaaacg tcgagtgaga gacaaggatg gagatctgaa 180
 gactcaaatt gaaaagctct ggacagaagt caatgccttg aagggaaattc aagccctgca 240
 gacagtctgt ctccgaggca ctaaaagttca caagaaaatgc taccttgctt cagaaggttt 300
 gaagcatttc catgaggcca atgaagactg catttccaaa ggaggaatcc tggttatccc 360
 caggaactcc gacgaaatca acgcctcca agactatggt aaaaggagcc tgccaggtgt 420
 caatgacttt tggctggca tcaatgacat ggtca 455

<210> 91
 <211> 488
 <212> DNA
 <213> Homo sapien

<400> 91
 actttgcttg ctcatatgca ttagtcaact ttataagtca ttgttatgta ttatattccg 60
 tagtagatg tgtaacctct tcaccttattt catggctgaa gtcacctctt ggttacagta 120
 gcttagcgtg gccgtgtgca tgcctttgc gcctgtgacc accaccccaa caaaccatcc 180
 agtacaaac catccagtgag agtttgc ggcaccagcc agcgttagcag ggtcggggaaa 240
 ggcacctgt cccactccta cgatacgcta ctataaagag aagacgaaat agtacataa 300
 tatattctat ttttatactc ttcttatttt ttagtgcacc tggttatgag atgctggttt 360
 tctacccaaac ggcctgcag ccagctcagc tccaggttca acccacagct acttggttt 420
 tggttttttt catattctaa aaccattcca ttccaaagca ctttcagtc aataggtgtaa 480
 gggaaatag 488

<210> 92
 <211> 420
 <212> DNA
 <213> Homo sapien

<220>
 <221> misc_feature
 <222> (1)...(420)
 <223> n = A,T,C or G

<400> 92

tctccggcag	gctctgcccc	ggtcgtagcn	agnnaaccta	taatcctgac	ctttttgt	60
gacaacctg	gtgctgagg	taactccatc	cattgttagt	gcctgtat	aatgggacg	120
attgcattt	tttcctgggt	gagcttcca	gaggctgaa	attttctccc	cacccttagt	180
ctgagatact	ttatcatgtat	cgancactc	cgtccactcc	acgtnttga	cccactcact	240
ggacaaagaa	acattgaaat	attcgccatg	ctctgtctgg	aacaatttga	atacccgggc	300
agcagcagag	cctcgatgnc	caggatattc	aatatggtct	tccactgaag	atgatggatt	360
tccttcaca	gntagaaaac	ttncnagggn	gtctaaatcc	aaggtgcagg	aangngngnc	420

<210> 93
<211> 241
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(241)
<223> n = A,T,C or G

<400> 93

accacgaatt	ncaacatcca	gatccaccac	tatcctaatt	ggattgttaac	tgngaactgt	60
gccccgctcc	tgaaagccga	ccaccatgca	accaacgggg	tggtgcacct	catcgataag	120
gtcatctcca	ccatcaccaa	caacatccag	cagatcattt	agatcganga	cacccttgag	180
acccttcggg	ctgctgnngc	tgcatacagg	ctcaacacga	tgcttgaagg	naacggncag	240
t						241

<210> 94
<211> 395
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(395)
<223> n = A,T,C or G

<400> 94

actctattnt	aattctgcct	ttttataactt	aattctaaat	ttttcccttc	taatttacaa	60
caaattttgt	gattttata	agaatctat	cctcccaat	tctcagattc	ttctctttc	120
tcctttat	cttgcctaa	attcagtata	agctttctt	gtattttagg	tttcatacgac	180
attttat	ctaaacacca	gcagttttc	agagacctaa	aatccagtat	aggaataact	240
gtgttagt	ttgaaaaagc	attaaagaca	ttttccctg	aaacatacag	aacatgtcat	300
gccaaatctc	ttgtttacat	aataaactgg	taataccggt	gaattgcaca	tacagat	360
atctccaaga	tagaataact	taaatattaa	aacgt			395

<210> 95
<211> 304
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(304)
<223> n = A,T,C or G

<400> 95

cgaggtacag	tgatngctcc	ccctggcaa	tacaatacaa	gaacngnggg	ttttgtcaaa	60
ttgaaacaag	gaaacagaac	cacagaaata	aatacattgg	ttaacatcag	attagttcag	120

gttactttt taaaatgtt aaagtacgag gggacttctg tattatgcta actcaagtt	180
actggaaatct cctgtttct ttttttttt taaatnggtt ttaattttt ttaattggat	240
ctatcttcctt ccttaacatt tcagttggag tatgttagcat ttagcaccac tggctnaaac	300
ctgt	304
<210> 96	
<211> 506	
<212> DNA	
<213> Homo sapien	
<400> 96	
acactgtcag cagggactgt aaacacagac agggtcaaag tgtttctct gaacacattt	60
agttggaatc actgtttaga acacacacac ttacttttc tggctctcac cactgctgat	120
attttctcta gaaaaataac ttttacaagt aacaaaaata aaaactctta taaatttcta	180
tttttatctg agttacagaa atgattactg aggaagatta ctcagtaatt tgttaaaaaa	240
gtaataaaaat tcaacaaaca ttgctgaat agctactata tgtcaagtgc tgtgcaaggt	300
attacactct gtaattgaat attattcctc aaaaaattgc acatagtaga acgctatctg	360
ggaagctatt ttttcagtt ttgatatttc tagcttatct acttccaaac taattttat	420
ttttgctgag actaatctt atcattttct ctaatatggc aaccattata accttaattt	480
attattaacc ataccctaag aagtac	506
<210> 97	
<211> 241	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(241)	
<223> n = A,T,C or G	
<400> 97	
attttctttt taattacttt agagagctag ggatgcaaatt gtttcagtt agaaaggcatt	60
tatttacttt tggaaattga acaagaaatg catctgtctt agaaaactgga gattatttga	120
tgttaggtaa aacatgttaat tgtntctctg gcaaatttgc atcantnatt ngaaaatgag	180
atattangaa aaaccaattt tccttaaattc tagncatct ttctttanaa gaacattana	240
t	241
<210> 98	
<211> 79	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(79)	
<223> n = A,T,C or G	
<400> 98	
ggcaaacana cttatgctgn ancngggttt tancaagggtt ttcaaagnaa aaancccatt	60
ngactttatg gaaaatatt	79
<210> 99	
<211> 316	
<212> DNA	
<213> Homo sapien	
<220>	

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<221> misc_feature
<222> (1)...(316)
<223> n = A,T,C or G

<400> 99
ccacatatgt aaaaccaga aagaccngnt tngcaacttc actgagagtt gagtcatctg      60
ggctgtcnac aggtgtctga cgtgtaaact tggaatcaaa ctgacttaca tcctcttcag     120
attgcAACAG aggttAAAG gggggctcca ccttcgagc cagaagtct tcccgatTTA     180
tgtgtctaaa gaatggatga gcttgaactt ctccagcgtc cccaggacca gctcccagac     240
gagaAGCAGC atttctttc agcagcttt taagcagatc tctggctct tgnGTgaggt     300
agggaggcaa attgag                                316

<210> 100
<211> 425
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(425)
<223> n = A,T,C or G

<400> 100
accgctttca gaaagtttat atgggttatt ctgcggcctc tctttatgc ctggcggcct      60
ctgtttatca accccaaacc aattacgtat ctggaaagtta tcaataccgt ggcacaggc     120
acttttgaca tttaattta ttacttttg ggaattaaat ctttagtcta catgttggca     180
gcatacTTAC ttggcctggg tttgcaccca atttctggac attttatagc tgtagcattac     240
atgttctaa agggncatga aacttactca tattatgggc ctctgaattt acttacctc     300
aatgtgggtt atcataatga acatcatgat ttccccaaaca ttcctggaaa aagtcttcca     360
ctggtgagga aaatagcagc tgaatactat gacaacctgc ctcactacaa ttctggata     420
aaagg                                425

<210> 101
<211> 156
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(156)
<223> n = A,T,C or G

<400> 101
actgacttgg gaatgtcaaa attctttatt atgatcttcc gagtggttgc ctgagctttg      60
ttggccctca actgcaggca gagaaccagg agcagggtgg cagggctggc cctgaacagg     120
agctggagca agcgcatgct ngagaaaaca gaaggc                                156

<210> 102
<211> 230
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(230)
<223> n = A,T,C or G

<400> 102

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actccaggcc gggnctcagg ttatcaaaag tgcaggagct ctgatcagca tggaccactt	60
cttccaaaga atttccctgc tgccgttg tagggttgt ggttaattcta taaccagtaa	120
tgtctgggt ggtgctcctc tcccaggaga ctgtgagcac tccagtgtca gggttgcct	180
ccagatgcaa gntngtnggt ggagacaatg gtgnaccac tttgtnnaca	230
<210> 103	
<211> 404	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(404)	
<223> n = A,T,C or G	
<400> 103	
actgtgaacc ctngggnttc nangcgacct acctggagct ggccagtgtc gtgaaggagc	60
agtatccggg catcgagatc gagtcgcgcc tcggggcac aggtgcctt gagatagaga	120
taaatggaca gctgggttgc tccaagctgg agaatgggg ctttcctat gagaagatc	180
tcatggagc catccgaaga gccagtaatg gagaaacctt agaaaagatc accaacagcc	240
gtcctccctg cgtcatcctg tgaactgcaca ggactctggg ttccctgcct gttctgggt	300
ccaaaccttg gtctccctt ggtcctgctg ggagctcccc ctgcctctt cccctactta	360
gctccttagc aaagagaccc tggcctccac tttgccttt gggt	404
<210> 104	
<211> 404	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(404)	
<223> n = A,T,C or G	
<400> 104	
accaggttat ataatagtat aacactgcc aaggacggat tatctcatct tcatacgtat	60
attccagtgt ttgtcacgtg gttgttgaat aaatgaataa agaatgagaa aaccagaagc	120
tctgatacat aatcataatg ataattattt caatgcacaa ctacgggtgg tgctgaacta	180
gaatctatat tttctgaaac tggctcctctt aggatctact aatgattaa atctaaaaga	240
tgaagtttgtt aaagcatcag aaaaaaaagt gggtattcct acaagtcaagg acattctacg	300
tgactataat ataatctcac agaaatttaa cattaatacn ttctaagatt taattcttag	360
antctngtta aacaaagtag ctccgtgga natgattggc atca	404
<210> 105	
<211> 325	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(325)	
<223> n = A,T,C or G	
<400> 105	
acagcagaag ccagtctang atgggtgtat tcaatttctg cctctagttat ttctttgtct	60
tgttttctt tcaattttaga agtgagcatt gtgttctcag ctatcagaac tttaagctgc	120
ccactatatt gagatgccct ttagctaat gattcctt tcaagtttag ggtcatctga	180
agttcagcat tctttctt taaaatctta atgtcctcaa agtatttatt ttccctttcc	240

tggtattggn gtttcagngt ggcttattcc agtttagca tggcaattnc cttaatcaac	300
atgcaatttt catgtaagag ataat	325
<210> 106	
<211> 444	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(444)	
<223> n = A,T,C or G	
<400> 106	
actgtcttca atnctatgcg tgcaagggtgtc taccacaggc aaacagttt ctccccattt	60
tgttagtaatg tgattttcctt attagcaaaa agaggtcacc agccccctgtta gacttaaggg	120
actcaagtca caggatgggg atttccttta aatattttttt atttngttgt ttgaactctt	180
gatgcaacat tgttagagcag ggtgttcagg acctgtgtg cccaaggac tgataaaagga	240
aaaagctcta tttattctttt ttgtgatttg atgcacagat gaaaaactta acacacaata	300
acagaagttt gncgttaata aatcacatcc taggcttca gcgcctncgt aagcagacga	360
catcttcagt tttctagtc ttgnagnntt aacacngnaa catcaatgtat gcatatgtnc	420
agaatcagt acaaagacca tccg	444
<210> 107	
<211> 287	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(287)	
<223> n = A,T,C or G	
<400> 107	
acctgcactc gnacntcagg cantaggcct ccacgtcatg gccaggcact ggcacatgggct	60
ccaccacgtg caggcagggtt cagtccttctt gggatacatt ctgggtgtaa atgtgccac	120
tgtatgtttctt ataaggtggg acagatgtat ttgcacccga tatcttcana actcttgg	180
gctncagctg ggggcaccaa caaacacccg accacagcca ccaaagataa nagcttcatg	240
cttatcangc ttgctgggccc agnnaagccg gacacctaca agccnc	287
<210> 108	
<211> 478	
<212> DNA	
<213> Homo sapien	
<400> 108	
acatgtgcaa gaatttggaa aagcagggca ttttcctca tctctccttag agggaatatc	60
acagcatctg tctctactgg tccacactgg actgcagaca atgtcaaaac tctggattt	120
gaatgcggct gatttcctt ccccttaag gagtttcca agaatttcat aaccatcagt	180
tgttatattt ccagcttctt tgatgtttt ttctataatt tcatacgat caatgtaaat	240
cttaacactt tttgaggtca ctacaatatg aaccttgta aaacttccat aaaataatgt	300
ctttacttct tctgtgtcaa atgtaacagt ttgcacctcg cctttgtat cttgttaaa	360
gaatgataac gtcttgctag aaggatctgc aatcactcca acttgtgggtt tgttagtctct	420
gtctgtgatt tgccaaattt gaaaagggtc actggagtt tctggagaa gtctgaat	478
<210> 109	
<211> 361	
<212> DNA	

<213> Homo sapien

<220>

<221> misc_feature

<222> (1)...(361)

<223> n = A,T,C or G

<400> 109

gaattttct tctanaataa gtattctgtt gacacagact attggtaaga tttcaacat	60
aaggtaatgc taggactggc ctccatgtcat gagttgttag taaagatctg gtctgttgg	120
tctccaaaag aagnntctta ctgttgtct ctcatgagtt ttctgttct gcttctctt	180
ttccatattg atatatacgg nttnnataat ggtnattgta attaaatatc tcctcatttt	240
tctcttttag gagatgtatgt tgcatttcc tctcaagaaa atgaatatca attgttatct	300
tgcttttgtt gncagcttc ttatgtgcat gaactaattt ctgttgaagc cacatatttt	360
t	361

<210> 110

<211> 305

<212> DNA

<213> Homo sapien

<220>

<221> misc_feature

<222> (1)...(305)

<223> n = A,T,C or G

<400> 110

acataatgac tnncanagtg aagctgattt gctgcgggttc tggagtaaat ataagcttc	60
cgttcctggg aatccgcact acttgagtca cgtgcctggc ctaccaaattt cttgc当地	120
ctatgtgcct tatcccaccc tnnaatctgn ctcctcattt ntcagctgtt ggatcagaca	180
atgacattcc tnttagatntg gcgatcaagg attccanacc tgngccaact gcaaacggtg	240
cctncaagga gaaaacgaag gcncacccaa atgnaaaaaa tgaangnccc ttgaatgtac	300
taaaa	305

<210> 111

<211> 371

<212> DNA

<213> Homo sapien

<220>

<221> misc_feature

<222> (1)...(371)

<223> n = A,T,C or G

<400> 111

cggggccag ccgggggtat tcagccatcg atcaaactca aaacctggaa tgatatccac	60
tctcttttc ttaagctcg gaaatattt caagtagaaag tccagaaatg catcgctaa	120
gatgcttcgg aatttgcattt catgcacata ggccttgaga aaactgtcaa actgatcctg	180
atcacccacc aagtggcca ggtatgagac aaagcagaaa ctttctcg agggggcttc	240
attataggtg tcgtccgggt caacgcctgg ttcaatctt acgcggagct tggtgagtgg	300
gttttcttcc caagtgtatgtt ccatgtgctg acgcagcaga ncccccccccc ttgcagcctc	360
caagcagggng t	371

<210> 112

<211> 460

<212> DNA

<213> Homo sapien

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<220>
<221> misc_feature
<222> (1)...(460)
<223> n = A,T,C or G

<400> 112
acatcttagg ttttnttcc tttantgtga agaggcgaaa ccaccaaccc acagctctgc      60
gtcgagttt tactagattt ctgcaaattt catggaaatct ttgctgttgt tcagtggtcc    120
atttatttgg a gccaaaaattt ctagggcgctt agaatggaa caaggttagtc agccaagcac   180
aaaaacataa caaaaacagga aacgccggac agaacagatg gatctagata gtagataatc   240
agaaaacacca aagaaaccac acccatgatg gcaggtggaa accaggctct ttctcatcgg   300
aggactttat cagccatcag catcaattt ccccatcctt gcagctgttc ttccagactt   360
gcagtctctg cagccagcag gttgggtgctt gctttttttt ccctccggca tcgtctcggg   420
gatgcagtctt ctacaagcgc aggccacccccc cccaaacgagt                         460

<210> 113
<211> 204
<212> DNA
<213> Homo sapien

<400> 113
gagaagacag cagagctgct ttccgcctct ttgagaccaa gatcacccaa gtcctgcact      60
tcaccaagga tgtcaaggcc gctgctaattt agatgcgcaa cttccctggtt cgagcctcct    120
ggccgccttag ctttggaaacctt gggaaaagaat atttgatcat gggtagat gggccaccc   180
atgacctcga gggacaccccc cagt                                         204

<210> 114
<211> 137
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(137)
<223> n = A,T,C or G

<400> 114
accgcaagaa atggacacgc aacgtcattt agacttttga catcgncgc tngacagtca      60
acgctgacgtt gggctatttttccctggaggt gtcccaagcc cctgaagaac cgtgatgtca    120
tcaccctccg ntccctg                                         137

<210> 115
<211> 278
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(278)
<223> n = A,T,C or G

<400> 115
gcggggcgctt ttntggactc gctcatttttac agagcatgcg tggctttcac ccttggcatg      60
ttctccgcgg gctctcgaa cctcaggcac atgcgaatgtt cccggagtgt ggacaacgtc    120
cagntcctgc cctttctcac cacggangtc aacaacctgg gctggctgan ttatggggct    180
ttgaagggag acgggatctt catcgtaanc aacacagtgg gtgctgcgt tcanaccctg    240
tatatcttttgcatatatctgc attactgccc tcggaaacg                         278

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<210> 116
<211> 178
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(178)
<223> n = A,T,C or G

<400> 116
.acaccgtcat angtcaaaaag tncagtgctg gccatcttgc atcaaatgtt cttaaaggcag      60
tgactggcta tcaaccacag nttctgtctc cccagntgca aacacaggat ccatgcaaca      120
gttctgagac catacactta gaaaccacng ggagatgcgg atcanatgca naactnnnc      178

<210> 117
<211> 360
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(360)
<223> n = A,T,C or G

<400> 117
actccccaat ggnnggattta ttactattaa agaaaccagg gaaaatatta attttaatat      60
tataacaacc tgaaaataat gaaaaagagg ttttgaatt ttttttttaa ataaacacct      120
tcttaagtgc atgagatggt ttgatggttt gctgcattaa aggtatttgg gcaaacaaaa      180
ttggagggca agtgactgca gtttgagaa tcagtttga ccttgatgat ttttgtttc      240
cactgtggaa ataaatgttt gtaataaagt gtaataaaaaa tcccttgca ttctttctgg      300
accttaaatg gtagaggaaa aggctcgta gccattgtt tctttgctg gttatagtt      360

<210> 118
<211> 125
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(125)
<223> n = A,T,C or G

<400> 118
gcgtcgtgct atgaccggac ttngtcttga aaggggatga cagcatggga ggcaatggnt      60
ncacatgtaa accccacact gaaagacaag gcactctctc cacagcagcc ccaacaacta      120
gcctc

<210> 119
<211> 490
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(490)
<223> n = A,T,C or G

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<pre> <400> 119 nacaaagaaa agcaaaaaga atttacgaag attgtgatct cttattaaat caattgttac 60 tgatcatgaa tgtagttag aaaatgttag gtttaactt aaanaaaatn gtattngat 120 tttcaatntt atgttgaat cngngtaata tcctgangtt ntttcccccc cagaagataa 180 agaggataga caacctcta aaatatttt acaatttaat gaaaaaaagn taaaaattct 240 caatacnaat caaacaattt aaatattta agaaaaaagg aaaagttagat agtatactg 300 aggtaaaaaa aaaattgatt caatttatg gtaaaggaaa cccatgcaat ttacctaga 360 cagccttaaa tatgtctgg tttccatctg ctgcatttc agacattta tttcccttt 420 actcaattga taccacaga aatatcaact tctggagtct attanatgtg ttgtcacctt 480 tctnaagctt 490 </pre> <pre> <210> 120 <211> 361 <212> DNA <213> Homo sapien </pre> <pre> <220> <221> misc_feature <222> (1)...(361) <223> n = A,T,C or G </pre> <pre> <400> 120 caggtacagt aaaattaaca cttccgttac agggaaatgt a tacgcaa atataaaaat 60 taaaaaggtga aaaaaagggtg acactggtt cctaagatac aatttactt ttacaaccag 120 ggccacagg tccaggctgc anagcggca tcagaagca gagcctncca cctgcttctg 180 ggggacctgg taataaaaat cagccatga tggcgtatg gcctctcaga caccacacgc 240 tgcctaaaca cctagagctc tggaaatagt caacaggaga gtgatttcca tggggaaat 300 tttaaanaag atgcacatgg gacaggcaat agaaagttt ccaaggntaa atttgtacc 360 t 361 </pre> <pre> <210> 121 <211> 405 <212> DNA <213> Homo sapien </pre> <pre> <220> <221> misc_feature <222> (1)...(405) <223> n = A,T,C or G </pre> <pre> <400> 121 acacaaaacc tttnacata ttggggcctt accgctccaa attgctactg atccttaag 60 ttcacaatat agaatttctt caccaattaa gtaataaccc tcattacaaa taaagtgcac 120 ctgataacca aactcgtaag tcccattgc agggactgct tggccattt aaggatcccg 180 tatatatgga catgttctc tataacaggc gtcatctgag acaggtagcc atgtatgatt 240 ccgatcacaa atatgtggg tggcaagagg aggtatata gagtatcctt tttacactt 300 ataatctact cgttcaccaa tctcatagta gggtttgg ttaaccatga gcctccatan 360 cttcaaatgt tgggtggctn ctcacaggca tcngcanaa ngagt 405 </pre> <pre> <210> 122 <211> 152 <212> DNA <213> Homo sapien </pre> <pre> <220> <221> misc_feature <222> (1)...(152) <223> n = A,T,C or G </pre>	
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<400> 122
 accccgctcc gttgnacacag atcgctgtct gcccactcca tcggccattc acttggcagg 60
 tgcgattggc agagccccgg agagtgtaac cgtcatagca gtggaaagag atctcatcac 120
 tcacattgtta gtagggagac cggggccaan ta 152

<210> 123
 <211> 336
 <212> DNA
 <213> Homo sapien

<400> 123
 acatctgaca tatttatata gcacataaat tagggagtgc tctgaccctt gcccgtggag 60
 cccaaagact gagcagggag gtgaacgcca gtccagaaag aaggtgtgg agccctgtct 120
 ctgtctcttc catcacgggg ctcccctagg gcctccccag gcctccttgg ctcagtccag 180
 gtgtctgcag gaggaaggtg ttgtctgcat tttagtgtctg agactgggtt tgaggaggca 240
 ccagataaaa ggagatacac ttgcagctat aaagttagct tcaaacccta gggcttgtaa 300
 ttccaagagg agggtgggaa ggcgaggcca tagtct 336

<210> 124
 <211> 253
 <212> DNA
 <213> Homo sapien

<220>
 <221> misc_feature
 <222> (1)...(253)
 <223> n = A,T,C or G

<400> 124
 ctgcaagagc ccagatcacc cattccgggt tcactccccg cctcccaag tcagcagtcc 60
 tagccccaaa ccagccaga gcagggtctc tctaaagggg acttgaggc ctgagcagga 120
 aagactggcc ctctagcttc taccccttgc ccctgttagcc tatacagttt agaatattta 180
 tttgttaatt ttattaaat gctttaaaaa aacaaaaaaaaa aaaaaaaaaaaa 240
 aaaaaaaaaaagntt gtn 253

<210> 125
 <211> 522
 <212> DNA
 <213> Homo sapien

<400> 125
 acaactgcaa gtctaagata atgttcattc attccatca taaatgtaac attctaaata 60
 ggtgtttct gatgtcatct gtcagaattt cttttaaact ttttcttcat cttcaacatt 120
 atcaaaagttc atccttattc ctcttgcctt gatttcggag agtttccaat ttttcaactt 180
 ttaaggcagc gattgtttt gcatctctgg tatttatctg ctcttcttgc aaatttctct 240
 ttgcttttc gtagaaataa aacttaacag ttggataggc cctgatccca gctttctggc 300
 atgtctgagc ataagcctga cagttactt ttccagctt cactttctt ttaatcatcc 360
 tagccaagag ctc当地attt ggagcaaaat tctggcaagg tccacacccaa ggagcataga 420
 aatcaatcac ccaatgattt ttcccttgc gaaacttttc actgaaagtc tgaggtgtta 480
 gatctgtgga tacttgaggt aaaaatccta gacccagat tc 522

<210> 126
 <211> 374
 <212> DNA
 <213> Homo sapien

<220>

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<221> misc_feature
<222> (1)...(374)
<223> n = A,T,C or G

<400> 126
ttttaagat attaactta cctttataaa tctttgtgt aaatgaaaaa aaaaatcaag      60
gcatacaaat ttcattgtgt tctacatttt taaataccat cctttgtctc cgtaaaaga    120
tttcatcca tttattcaaa aacctttaa gttcaactgt ccaatttaag acagagtcaa    180
gacattttg agtatctgaa ctaaggcattg tcttgactga aacgaagtaa gaactcaatg   240
agagtcccttgggcctccc aggcatgcct ttccgttagat agggaaacttc atctttgtt     300
gnccatcacgc ctgctatgtc taaatgtgcc cacttaggat gagttacgaa ttcttcagg   360
aatgtcgac ctgt                                374

<210> 127
<211> 130
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(130)
<223> n = A,T,C or G

<400> 127
aaagccaaga cngccattgg cactgctatg gtaaggncac agggcancca gggccttctg      60
gcaaaaggng atacnaccag cactatnaac agacaggaca tggttgagag gnagnctaca    120
caantcctaa                                130

<210> 128
<211> 350
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(350)
<223> n = A,T,C or G

<400> 128
acactgattt ccgntnaaaa gaancatcat ctttaccttgc acttttcagg gaattactga     60
actttttctt cagaagatag ggccacagccat ttgccttggc ctcacttgc gggctctgc      120
ttgggtcctc tggctctttt ccaagnttcc cagccactcg agggagaaat atcggggaggt    180
ttgacttccct ccggggctt cccgagggct tcaccgtgag ccctgcggcc ctcagggctg   240
caatccttggat ttcaatgtct gaaacctcgat tctctgccttgc ctggacttctt gaggccgtca 300
ctgcccactctt gtcctccagc tctgacagct cctcatctgt ggcctgttga                350

<210> 129
<211> 505
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(505)
<223> n = A,T,C or G

<400> 129
acaataccaa agcttcataaa tgctaaagaa aacccaaaaca aaagacaatg gtttacacag      60

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ggaaaataacc ctaaggcaat atgaaaacag tcataattta ttactgataa agagtaaagg	120
catccccc atagaggggg ggaattcaca gggAACACTA attatatcg atgaaccacg	180
gggatagaaa ataggccat tttaaaattt cattgagaaa ttattactt ttctccacaa	240
ctgtgattct atacaaaata taaaccctgc aaaccttatg tgctacctga cagataaaag	300
tagcaggagc cagactttg aagacttga gactgattc tacaaagtcc aggaagagca	360
atgattccag tgtcagtc tgatgcattgt gtgagcctaa catgttattc agctctggtt	420
gcagccccat ctacatgggg cccagttgt ttttagggag tcacagatta ngcaggcaac	480
cgaggggcat gattaaaaa gcaca	505
<210> 130	
<211> 526	
<212> DNA	
<213> Homo sapien	
<400> 130	
acaaaagagc ctgattctt ttaattccac aaataacctag catctcaaag taacatgtaa	60
acaaacttct atgctgctca atgaatcctt ccaattcga taataaaacta aatagtattg	120
gatctagttat atgactttca tgttaagtt atggttctat ccattactt aacaatatta	180
ctgatgtaac agagaaaaat tttcaactat tgtacttatt taaaacaaac tgacaagttc	240
aagcacctgt cttcagaaaa gccagcagca tttttttttt ttaacatac tcaaagtaag	300
atttggctta agcccttaat accttctga acagccatgc aactaaacac cctcaggaga	360
tgttacataa gggagagaag aacatggagc aatttgcact tttccctta gataatatta	420
acaaggtaaa gcaaatccag atcttatga atgaatggct gtcatgttta atacacttgg	480
agctctataa aactagagcc actatcatat atgttataat agatat	526
<210> 131	
<211> 477	
<212> DNA	
<213> Homo sapien	
<400> 131	
ctcagtttc ccagcaacag atgctcctga gcaatttatt agtcaagtga cggtgctgaa	60
atactttct cattacatgg aggagaacct catggatggt ggagatctgc ctatgtttac	120
tgtatattcga agacctcgcc tctacctcct tcagtggcta aatctgata agggccctaat	180
gatgtctttt aatgtatggca ctttcaggt gaatttctac catgtatata caaaaatcat	240
catctgttagc caaatgaag aataccttct cacctacatc aatgaggata ggatatctac	300
aacttcagg ctgacaactc tgctgatgtc tggctgttca tcagaattaa aaaattgaat	360
ggaatatgcc ctgaacatgc ttttacaaag atgtaactga aagactttc gaatggaccc	420
tatggactc ctctttcca ctgtgagatc tacagggAAC ccaaaaagaat gatctag	477
<210> 132	
<211> 404	
<212> DNA	
<213> Homo sapien	
<220>	
<221> misc_feature	
<222> (1)...(404)	
<223> n = A,T,C or G	
<400> 132	
accacacgan cgggnatcnt ttgnacatag tgagacccgg ctgattccca tacatgaatc	60
cattcatgga gtgcattttt ttagatncct gaaagtcttc atcttcctta tccacctgtat	120
cagggcagt tgttaacatn cctaatattt tcttccagga gtaaaactctc attctcatca	180
aatactgtat gaaacaaata gaattccctt tctacatctt tctgtctccc atttgcatat	240
aaacttcctt tttgcataat tttcattggc ccaataagcc cagtgatata atctttatgt	300
ggatccacag cagaataata catcttagct agacacacag gatctgcatt tacgngggtc	360
ctacttcttt ggggacagcc cttcatacgn gaatgttnt gtgg	404

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<210> 133
<211> 552
<212> DNA
<213> Homo sapien

<220>
<221> misc_feature
<222> (1)...(552)
<223> n = A,T,C or G

<400> 133
accccaaatt atctctctcc tgaagtcctc aacaaacaag gacatggctg tgaatcagac 60
atttgggccc tgggctgtgt aatgtataca atgttactag ggaggcccccc atttgaact 120
acaaatctca aagaaaactta taggtgcata agggaaagcaa ggtataacaat gccgtcctca 180
ttgctgctc ctgccaagca cttattgtc agtatgttgt caaaaaaccc agaggatcg 240
cccagttgg atgacatcat tcgacatgac tttttttgc aggcttcac tccggacaga 300
ctgtcttcta gctgttgtca tacagttcca gatttccact tatcaagccc agctaagaat 360
ttctttaaga aagcagctgc tgctttttt ggtggcaaaa aagacaaagc aagatatatt 420
gacacacata atagagtgtc taaagaagat gaagacatct acaagcttag gcatgattt 480
aaaaagactt caataactca gcaacccagc aaacacaggg acagatgang agctccacca 540
cctaccacca ca 552

<210> 134
<211> 496
<212> DNA
<213> Homo sapien

<400> 134
acattgatgg gctggagagc agggtggcag cctgttctgc acagaaccaa gaattacaga 60
aaaaagtcca ggagctggag aggcacaaca tctccttggt agctcagctc cgccagctgc 120
agacgcta at tgctcaaact tccaaacaaag ctgcccagac cagcacttgtt gtttggattc 180
ttctttttc cctggctctc atcatcctgc ccagcttcag tccattccag agtcgaccag 240
aagctgggtc tgaggattac cagcctcagc gagtgacttc cagaaatatc ctgaccacaca 300
aggacgtAAC agaaaatctg gagacccaaag tggtagagtc cagactgacg gagccacctg 360
gagccaagga tgcaaattggc tcaacaagga cactgcttga gaagatggg gggaaGCCAA 420
gaccCAGTGG ggcgcattccgg tccgtgctgc atgcagatga gatgtgagct ggaacagacc 480
tttctggc cacttt 496

<210> 135
<211> 560
<212> DNA
<213> Homo sapien

<400> 135
actgggagtg atcactaaca ccatacgtaat gtctaattt cacaggcaga tctgcttggg 60
gaagctagtt atgtgaaagg caaatagatc catacagtag ctcaaaaggc aaccataatt 120
ctctttggc caggtcttgg gagcgtgatc tagattacac tgccaccattc ccaagttat 180
ccctgtaaaa cttaactctca actggagcaa atgaactttg gtcccaataa tccatcttt 240
cagtagcggtt aattatgctc tgtttccaaac tgcattttt ttccaaattga attaaagtgt 300
ggcctcggtt ttagtcattt aaaattgttt tctaagtaat tgctgcctctt attatggcac 360
ttcaattttt cactgtctt tgagattcaa gaaaaatttc tattttttt tttgcatcca 420
atgtgcctg aacttttaaa atatgtaaat gctgccatgt tccaaacccca tcgtcaagtg 480
tgtgtgttta gagctgtgca ccctagaaac aacatattgc ccatgagcag gtgcctgaac 540
acagacccct ttgcatttcac 560

<210> 136
<211> 424

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<212> DNA
 <213> Homo sapien

 <220>
 <221> misc_feature
 <222> (1)...(424)
 <223> n = A,T,C or G

 <400> 136
 accagcaaat ctccatttagc atttctcagg tttcatgatc ctttcagat atgttggttg 60
 attttatgtatataatgc tt agaaaacaaaa atccacactga tattaaaaca aaccaaaaaaa 120
 aatcataaaaa gcaagcaa at gaacaaaaaa ccctagttt gttgtgctt tcttcacat 180
 ttccctacagg gagatttgc tatctcagat acttcaaaa tctaatacggt aagtaaaatt 240
 agtgccttaa ccaaacagta agataccaa gaatcccca tcacaagttt ctgaatcaaa 300
 cttctcatga catttgcggt atattcagat ttgaagattt tttaaatttta gaatttaaaa 360
 caaacttttag actgctgatt ttccatattt caaagactgt agctgtntgc agcatataaa 420
 tgga 424

 <210> 137
 <211> 392
 <212> DNA
 <213> Homo sapien

 <220>
 <221> misc_feature
 <222> (1)...(392)
 <223> n = A,T,C or G

 <400> 137
 tgcggggntg aaggcttagca aaccgagcga tcatgtcgca caaacaaaatt tactattcgg 60
 acaaatacga cgacgaggag tttgagtatc gacatgtcat gctgcccaag gacatagcca 120
 agctgggccc taaaaccat ctgatgtctg aatctgaatg gaggaaatctt ggcgatcagc 180
 anagtcaggg atgggtccat tatatgatcc atgaaccaga acctcacatc ttgctgttcc 240
 ggcgcccact acccaagaaa ccaaagaaat gaagctggca agctactttt cancctcaag 300
 ctttacacag ctgnccctac ttcttaacat ctttctgata acattattat gctgccttcc 360
 tgttctcaact ctganatnta aaagatgttc aa 392

 <210> 138
 <211> 284
 <212> DNA
 <213> Homo sapiens

 <220>
 <221> misc_feature
 <222> (1)...(284)
 <223> n = A,T,C or G

 <400> 138
 tgcctgtgca cctctttgct tgaaatatgg caagacttgg aaaaatgttt gcccttagaa 60
 tctatctcac tacttttagtt agttgtctcc tttgggcctg ggcacagttc tggccctgat 120
 ctggaacaga ctcccttttc taaaactgaa cttgaccaca tcaaaagntt gnaaaacaat 180
 ctccatggta attaaacttg cattcaacac catatggnaa cagaagatgg caggaggata 240
 anatncagat cttatgatct ttccangnan ggcatgttac atga 284

 <210> 139
 <211> 249
 <212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(249)

<223> n = A,T,C or G

<400> 139

gaggaagggg ggactgaatc tancaccntg acngaactag agacagccat gggcatgatc 60
atagacnnct ttacccgata ntcgggcagc gagggcagca cgcagacccct gaccaagggg 120
gagctcaagg ggctgtatgga gaaggagcta ccaggcttcc ngcagagnng aaaanacaag 180
gangccgtgg ataaaattgtt caaggaccta gacgcnatg gaggatgccccc aggtggactc 240
cagcgagt 249

<210> 140

<211> 390

<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(390)

<223> n = A,T,C or G

<400> 140

tcataatggc tggggcagct ataatnnact acaanaatca natgtttcac atctagacct 60
cgggcagcaa cagaggtgc cacaagaatg ttgcangtcc cattcttaaa gtcatttatg 120
atgctatctc tgtcatattg atcaatgcct ccatgaagag acatgcaagg ataagatgct 180
ctcattaaat ccttaagaag accatcagca tgttcctgct tatccacaaa tataatgaca 240
gatcctgact cttgataatg gcctagaagc tcaagtaact tcaagaattt ctttcttct 300
tcaatcacaa tcacttgtn gctccacatct gagcaaacca cactcctgcc tccaacttgt 360
acctcccccg ggcggcgct caaggcgaa 390

<210> 141

<211> 420

<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(420)

<223> n = A,T,C or G

<400> 141

gacactcagg gaaaagcatn ngncaaanag agcttaaat gcatcgccaa cggggtcacc 60
tccaaggctc tcctcgccat tcggaggtgc tccacttcc aaaggatgat tgctgaggtg 120
caggaagagt gctacagcaa gctgaatgtg cgcanccatcg ccaagcggaa cccngaagcc 180
atcaactgagg tcgtgcagct gcccacatcac ttctccaaaca nataactataa cagacttgn 240
cgaaggctgc tgaaatgn gtaanacaca gggcagcaca atcaggagac agcctgatgg 300
anaaaantgg gcctancatg gccaggcctc ttccacatcc tngcangaca gaccactgtg 360
cccaaacaca cccnctgagc tgacttnac aggagacgca cnaaggagcc cggcagangc 420

<210> 142

<211> 371

<212> DNA

<213> Homo sapiens

<400> 142

gggttcgaca atgctgatcc gcaattagaa gacactggta agctgtgtt cactggcctt 60
cattgaaatc ttcaaggata tagccagctc ctgctcgaa ctgggattct gtatactgct 120
tgttgaaagg aggaatttcc aaaaatttcc ccttcttc actgcttcct gtaggaccat 180
ctggcagtt ggagcggctg gccaaacttgt cactggttgt ggccatggta aggagaaatg 240
cgtagcccag aaacaaggc ttgtttagag gcaaaggccc tctctgtct tccagggcag 300
agggttcacc ggtgttgtct ccactctcac aggggctcac aaactctcct gcccctactt 360
gcaccaggtt t 371

<210> 143
<211> 270
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(270)
<223> n = A,T,C or G

<400> 143
ggtggctgtg atnaccttn ttagttaca aataaaaaag ntaaaaagaa atactgtgtt 60
tagggttaagg taacanntc atctaattcag aggagagtga agangaggcn ctgccttcta 120
ggngctgtga ctttcctt ttctgngattc ttcnccacat tggnaacat cttccccgt 180
atgctggaan tacttcggng ttctgcggtg gccatgntga acatctgatg aactgaaant 240
ncatccnaat gcacacgaag anatagnncna 270

<210> 144
<211> 259
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(259)
<223> n = A,T,C or G

<400> 144
ttctctttgc ttttataat tttaaagnaa ataacacatt taactgtatt taagtctgtg 60
caaataatcc ttcagaagaa atatccaaga ttctgtttgc agaggtcatt ttgtctctca 120
aagatgatta aatgagtttgc ttccagata aagtgcctt gtccagnaga actcaaaagg 180
ccttcaagct gttcagtaag tgttaggttca gataagactc cgncatacga attccagctt 240
ccctgtccca ctgtacctc 259

<210> 145
<211> 433
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(433)
<223> n = A,T,C or G

<400> 145
accacatnta ccatagtgtt attagtttta atttcacat gaatcaaagg tttcccttca 60
tgtctattta cagtccaaatt gtgccaaact cttacttgtt tgctgactaa caaggcattt 120
agggtgtgcag catccttagag tgctccaggg cagtgtcagc gttctcggtt gtaaaagggtg 180
ccacttggta gcaatgatat tccagaattt aatgggttt tggtccatg gagactgcat 240
ttatataaat gtagcctgtt gcttaagtta actaaacctt atgctgtgt taaaaacagt 300

ttatTTTaat attaaaatac agttgattag caacagcggt gctgtattt aagagacact 360
ttattGGAAG tgcaatcata gttattgtt ttcacaattt tacagngcat tctaattact 420
gatgggtgca att 433

<210> 146
<211> 576
<212> DNA
<213> Homo sapiens

<400> 146
acctcaggcc tgtgcaccc tttgcttgaa atatggcaag acttggaaaa atgtttgcc 60
ttagaatcta tctcaactt ttagtttagt gtctcccttg ggctctggca cagttctggc 120
cctgatctgg aacagactcc ctttctaaa actggaccc tt gaccacatca aaagtttga 180
aaacaatctc catggtaatt aaacttgcatt tcaacaccat atggtaacag aagatggcaa 240
aggataagat tcagatctta gatcttcca agtagggcat gttagatgat agaaggatta 300
gttgcaagct ggatctgagc tcaggcttg ggcatgaagga aactgtctcc catgtggttt 360
ggaagagttt ggggctccct gagctctatt gtgaactata cgggtttcat ccaaggaatg 420
gtatgatgtg ggcataaaaac cattcttcag acaactgaag atggcccct tctgttagcca 480
gaaacactag ctgtcctgca ttgccatttc cttaaaaaa ggcggcctgc agaaggaaag 540
gccataatta attaaaaggc ttaatgaagt tttgga 576

<210> 147
<211> 300
<212> DNA
<213> Homo sapiens

<400> 147
ccagccccca ggaggaaggt gggctctgaat ctagcaccat gacggaacta gagacagcca 60
tgggcatgtat catagacgtc tttacccgat attcggcag cgagggcagc acgcagaccc 120
tgaccaaggg ggagctcaag gtgcttatgg agaaaggagc taccaggctt ctgcagagtg 180
gaaaagacaa ggatgccgtg gataaattgc tcaaggacct agacgccaat ggagatgccc 240
aggtggactt cagtgagttc atcgtgttc tggctgcaat cacgtctgcc tgtcacaagt 300

<210> 148
<211> 371
<212> DNA
<213> Homo sapiens

<400> 148
acataatcct cataatggtt gggcagcta taatttacta caagaatcag atgtttcaca 60
tctagaccc tggcagcaac agaggttagcc acaagaagtt tgcaggtccc attcttaaag 120
tcattttatga tgctatctct gtcatttattga tcaaattggcc tccatgaaga gacatgcaag 180
gataagatgc tctcattaaa tccttaagaa gaccatcagc atgttcctgc ttatccacaa 240
atataatgac agatcctgac tcttgataat ggcctagaag ctcaagtaac ttcaagaatt 300
tctttcttc ttcaatcaca atcacttgc tgcctcacatc tgagcaaacc acactcctgc 360
ctccaacttg t 371

<210> 149
<211> 585
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(585)
<223> n=A,T,C or G

<400> 149
 cgaggtacan cactgctaaa tttgacactn angaaaaagc attcgtaaa gagagcttaa 60
 aatgcacatgc caacggggtc acctccaagg tcttcctcgc cattcgagg tgctccactt 120
 tccaaaggat gattgcttag gtgcaggaag agtgctacag caagctgaat gtgtgcagca 180
 tcgccaagcg gaaccctgaa gccatcactg aggtcgtcca gctgccaaat cacttctcca 240
 acagatacta taacagactt gtccgaagcc tgctggaatg tgatgaagac acagtca 300
 caatcagaga cagcctgatg gagaaaaattg ggccta 360
 tgcagacaga ccactgtgcc caaacacacc cacgagctga ctcaacagg agacgcacca 420
 atgagccgca gaagctgaaa gtcctcctca ggaacctccg aggtgaggag gactctccct 480
 cccacatcaa acgcacatcc catgagagtg cataaccagg gagaggnat tcacaacctc 540
 ccaaactagt atcatttttag gggngttga cacaccagtt ttgag 585

<210> 150
 <211> 642
 <212> DNA
 <213> Homo sapiens

<220>
 <221> misc_feature
 <222> (1)...(642)
 <223> n=A,T,C or G

<400> 150
 acttnccgggt tcgacaatgc tgatccgcaa ttagaagaca ctggtaagct gtgttacact 60
 gggcttcatt gaaatcttca aggatatgc cagtcctcgc tcaagctgg gattctgtat 120
 actgcttgtt gaaaggagga atttccaaaa attcctcctc ttcttcactg ctccctgttag 180
 gaccatctgg cagtttggag cggctggcca acttgcact gtttggcc atggtaagga 240
 gaaatgcgtt gcccagaaac aaggcttgc tgagaggcaa aggccctctc tgctcttcca 300
 gggcagaggg ttcaccgggt ttgtctccac tctcacaggg gctcacaaac tctccgtccc 360
 ctactgcacc agttttact gtggcagact tgcgacccctcg cttggcaggg gaccgttcc 420
 cttcagaagt gataagttt ctttgcctg agagaactcc catggaggca cgaggacttt 480
 ctgtgatctt tccggtaggg gttgtgtgc tactggaggc agtangggtg gctggggagc 540
 tgacgttact ggcgcgtttc cgcttccttc caccaattt ctaagctgat atctgctgcc 600
 tttgttaagaa gnggtactgc ttcatanggg ccaagcccat ac 642

<210> 151
 <211> 322
 <212> DNA
 <213> Homo sapiens

<220>
 <221> misc_feature
 <222> (1)...(322)
 <223> n=A,T,C or G

<400> 151
 ntggacaaac atcttccccg ctagctgg attacttcgg tttctgcgg tggccatgg 60
 gaacatctga tgaactgaaa ttccatcgga atgcacagga agatata 120
 aatgtccctt ccaggaccac catactgggg aagttcttc gggtgcctgc naatgggctg 180
 caccctgggg ctggggcccg a gctctagctc tgtcatgcca tggccactga aatcggttt 240
 cagatgatta gtctcttcat gcccgtcca ttttcgggtt ttctccagt gttcagaaat 300
 tcaaatgatt aacttctggg aa 322

<210> 152

<211> 262
<212> DNA
<213> Homo sapiens

<400> 152
acaaagtctt ctcttgctt tttataattt taaagcaaat aacacattta actgtattta 60
agtctgtgca aataatcctt cagaagaaat atccaagatt ctgtttgcag aggtcatttt 120
gtctctcaa gatgattaaa tgagttgtc tttagaataa agtgctcctg tccagcagaa 180
ctcaaaaggc cttcaagctg ttcaagtaat gtatgttcaga taagactccg tcatacgaat 240
tccagttcc cgtccccact gt 262

<210> 153
<211> 284
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)
<223> n=A,T,C or G

<400> 153
ctcgggagta aaaggtagcca cttggtagca atgatattcc agaattaaat gggttttgt 60
tgccatggag actgcattta tataaatgtt gcctgttagt taagttaact aaacctaaatg 120
ctgctgttaa aaacagttt ttttaatatt aaaatacagt tgatttagcaa cagcggtgct 180
gtatttaag agacactttt ttggaaagtgc aatcatagtt atttggtttc acaattttac 240
ngtgcattct aattactgtat gggngcaatt acttttaatc gnng 284

<210> 154
<211> 531
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(531)
<223> n=A,T,C or G

<400> 154
acccacccta aatttgaact cttatcaaga ggctgatgaa tctgaccatc aaataggata 60
ggatggacct ttttttagt tcattgtata aacaaatttt ctgatttggc cttattccc 120
aaaggattag gtctactctt gtcattcac tcttcaaag ctctgtccac tctaactttt 180
ctccagtgtc atagataggg aattgctcac tgcgtgccta gtctttcttc acttacctgg 240
cctctgatag aaacagttgc ccctcttatt tcataaggta gaggacttgt gaccctggat 300
ggttctaaat gaaaaaaagca ccgccagatt gtgaaacctg gcttcaacat cagcattctg 360
aaaatattca tcaccatgtat gtctgagagt gttcgatga tgctgaacaa atgggaggaa 420
cacattgccc aaaactcacg tctggagctc tttcaacatg tctccctgtat gaccctggac 480
agcatcatga agtgtgcctt cagccaccag ggcagcatcc agtngacag t 531

<210> 155
<211> 353
<212> DNA
<213> Homo sapiens

<220>

<221> misc_feature
<222> (1)...(353)
<223> n=A,T,C or G

<400> 155
tcttgacaag actgagagag ttacatgttg ggaaaaaaaaa agaagcatta acttagtaga 60
actgaaccag gaggcattaag ttctgaaatt ttgaatcatc tctgaaatga agcagggta 120
gcctgccctc tcataatcc gtctgggtgc cagaactcaa ggtagtgg acacatcccc 180
ctgttagaga ccctcatggg ctaggacttt tcataatggg tagattcaag acctttac 240
canaattatg taaaactgtga ttgtgtttt gaaaaattat tatttctaa aaccatttaa 300
gtcttgtat atgtgtaaat gatcacaaaa atgtatTTTaaatgttc tgt 353

<210> 156
<211> 169
<212> DNA
<213> Homo sapiens

<400> 156
agtttgttct actacatttg tggccacta gttcacccat ctgtgttgat aagcggttacc 60
accatttgc ctttctatacg cctctttac aatgttgctc acttcatcaa caacaaaagc 120
agtctccccc gcagcctggt agtctccat cttccctccg ggcgtccc 169

<210> 157
<211> 402
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(402)
<223> n=A,T,C or G

<400> 157
gttaactacc cgctccgaga cgggatttat gacgaggcct atgaggccat tttcaagccg 60
gtcatgtcca aagtaatgaa gatgttccag cctagtgcgg tggcttaca gtgtggctca 120
gactccctat ctggggatcg gttaggntgc ttaatctac tatcaaaggaa cacgccaagt 180
gtgtggaaatt tgtcaagagc ttaacctgc ctatgctgat gctggggagggc ggtggttaca 240
ccattcgtaa cgttgcggc tgctggacat atgagacagc tggccctgt gatacggaga 300
tccctaatacg gcttccatacg aatgactact ttgaataactt tggaccagat ttcaagctcc 360
acatcgtcc ttccaaacatcg actaaccaga acacgaatgaa gt 402

<210> 158
<211> 546
<212> DNA
<213> Homo sapiens

<400> 158
actttgggtt ccagacttca ctgtccttag gcattgaaac catcacctgg tttgcattct 60
tcataactga ggttaactta aaacaaaaat ggttaggaaag ctttcctatg ctgcggtaa 120
gagacaaatt tgctttgtt gaattgggtgg ctgagaaagg cagacaggcc ctgattaaag 180
aagacatttgc tcaccactag ccaccaagtt aagttgtggaa acccaaagggt gacggccatg 240
gaaacgtaga tcatacgctc tgctaaatggaa ttaggggaaatggaaatggaaatggaaatggaa 300
ccaaatggat cctgtggta cagtgaaatgaa ccactcctgc tttatTTTtc ctgagattgc 360
cgagaataac atggcactta tactgtatggg cagatgacca gatgaacatc atcatccaa 420
gaatatggaa ccaccgtgttcaata gatTTTccc tggatgttag gcattcctgc 480
catccattgg cacttggctc agcacagttt ggcacaaacaag gacataatag acaagtccaa 540

aacagt 546

<210> 159
<211> 145
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(145)
<223> n=A,T,C or G

<400> 159
acttttgcta taagtttcctt aaaaatattt aatactttt ttttcaatt taaattaaat 60
ctnttgatga acaggggggg gntggcaaaa ttccaagcn ctggactgga attttganan 120
aggcatttac ngaccctnat aactt 145

<210> 160
<211> 405
<212> DNA
<213> Homo sapiens

<400> 160
tgtaaatcgc tgtttgatt tcctgatttt ataacaggc ggctggtaa tatctcacac 60
agttaaaaaa atcagccctt aatttctcca tgtttacact tcaatctgca ggcttcttaa 120
agtgacagta tcccttaacc tgccaccagt gtccccctc cggcccccgt cttgtaaaaa 180
ggggaggaga attagccaaa cactgtaagc tttaagaaa aacaaagttt taaacgaaat 240
actgctctgt ccagaggctt taaaactggt gcaattacag caaaaaggaa ttctgttagct 300
ttaacttgta aaccacatct ttttgcact tttttataa gcaaaaacgt gccgtttaaa 360
ccactggatc tatctaaatg ccgatttgag ttcgcgacac tatgt 405

<210> 161
<211> 443
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(443)
<223> n=A,T,C or G

<400> 161
tttgctttta atgaaggaca agggattaag acncatagag actggccana caaatggaa 60
accgaccaga ccagcccatg accaaaatat cacaggcaga ccacccacaa atgcagaggc 120
ctcagagtcc acagtggcg gttggAACCC agggccccag ggaatcttc agctgcattc 180
cggctgtat cggcgggcaa caggttagagg tgctggaggg ggctgagtcg tgatttcgg 240
tgtctgtcat attcgatcaa gtgtgtcata gagcttcctg tttcatctcc cagttattca 300
aggagaggct ggtggctcca cttcccccagg aactgtgctg tgaagatctg aagacaggca 360
cgggctcagg caccgcttgt ctggaatgtc aatttgaaac taaaaagca gcgaccatcc 420
agtcatat tttccat tcc 443

<210> 162
<211> 228
<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(228)

<223> n=A,T,C or G

<400> 162

tcgttatcaa aatggaagac accaaaccat tactggcttc taagctgaca gaaaaggagg 60
aagaaatcggt gacttagtgg agtaaatttt atgctnctc agggaaacat gaaaaatgcg 120
gacagtataat tcagaaaggc tattccnagc tcaagatata tnattgtgaa ctanaaaata 180
tagcanaatt tgagggcctg acagacttct canatacnntt caagttgt . 228

<210> 163

<211> 580

<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(580)

<223> n=A,T,C or G

<400> 163

acccaaggct acacatcctt ctgtgaaaca gtctcacgga gactctcaga atcccaagaa 60
ttttcttcaa ccttcttttgg tttgattct gaagggaca tctgatctgc tctcaatgtt 120
tgttcattct tcaattccaa ggctttatTTT ggaacagact ttgcatttca atggcaggct 180
cgaaggcaga tggcttcctcg ggaggctctg ctttggaaagt ttgcntgtcc atcaattcta 240
aggctttagt tggaaatagaa actttcatTC tgcaggggac cttcagaaaa ccattcattat 300
caggagactc ttctaatTTT ccatttattt tatctatttcc tttttgatgc gcagccttgg 360
gtanacacac atccttctgt gaaacagtct cacagagact ctcagaatcc caagaacttt 420
cttcatagtc cttttgtttg gattctgatg ggagtatctc atctgctctc aatgtttgtt 480
catttttcaa ttccaaggct ttatTTGAA cagacttttgc catttcaatg gcaggctcga 540
aggcagatgg cttctcgaaa ggctctgctt tgaaaagttg 580

210> 164

<211> 140

<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(140)

<223> n=A,T,C or G

<400> 164

acttataatct tttgggncttgg ggcttctcaa agttcacgac agacataggc actctcacag 60
tatcaagccc atttaccgnc acctcacacc aataactcgcc ccaccgngng ataggnctcg 120
ctggnaactt taatgnatgn 140

<210> 165

<211> 370

<212> DNA

<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(370)
<223> n=A,T,C or G

<400> 165
acatggagcc actgccacca gtggtgatgg aaagcactgc cttcttactc cggaagggtc 60
ctttgtcata catggcagcg taagtgtaaag caaactctcc tatgaacact cgctcaaacc 120
agccttcag aatggcaggg actccaaacc actgcnnnnn ggaactggaa tatcacaagg 180
tctgggctt ccagttctt ttgttcagcc acaatatctg ggctcanatg gncttctta 240
taagccagaa cagactcggn aggatactga aagttcgcag ggnccctcan tttacctgng 300
atgncccttn tggaaatgtat gggattgaag ntcatggnat aaaggncgca ctnaccacc 360
tccattcttt 370

<210> 166
<211> 258
<212> DNA
<213> Homo sapiens

<400> 166
gtcaaaaatgc atgattttta tcttagttct tcattactgc attgaaaagg aaaacctgtc 60
tgagaaaaatg cctgacagtt taatttaaaa ctatgggtta agtctttgac aagaaaaaaaaa 120
aacaacaaaa cacttcttc catcagtaac actggcaatc ttccctgttaa ccactctcct 180
tagggatggt atctgaaaca acaatggtca ccctcttgag attcggttta agtgtaattc 240
cataatgagc agagggtgt 258

<210> 167
<211> 345
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(345)
<223> n=A,T,C or G

<400> 167
ggtcagccaa acacccagga tctctgtaaa actgaagaac aggncaatgc caccacaaaa 60
tctcaaaacc tctccagcat atttccttat gattggagca catggngagc acnancggc 120
acttttaaca canctagcca gacagggngnc atttgggtta acacttcgga acccacagca 180
ntttanantt ctctggatgt catttcgagc acttgtattt attggtcann tttctgtatc 240
tngcgttgg ttagccctga accaggagca acagggnncag cttctggagg ntgggtggaa 300
caatacggca agtgnntngaa atgacatcca acctncngaa atgac 345

<210> 168
<211> 61
<212> DNA
<213> Homo sapiens

<400> 168
gatagtgtgg tttatggact gaggtcaaaa tctaagaagt ttgcgcagacc tgacatccag 60
t 61

<210> 169

<211> 344
<212> DNA
<213> Homo sapiens

<400> 169
acattggtgc tataaatata aatgctactt atgaagcatg aaattaagct tctttttct 60
tcaagtttt tctcttgct agcaatctgt taggcttctg aaccaagacc aaatgttac 120
gttcctctgc tgcataccaa cgtaactcca aacaataaaa aatctatcat ttctgctctg 180
tgctgaggaa tggaaaatga aaccccccacc ccctgaccacc taggactata cagtggaaac 240
tgttcattgc tgatgaatgc agcagtccacc aaaaaataca cccaatcttc cagataacct 300
cagtgcactt taggaaatca aaaattacct ggaagcaatt tagt 344

<210> 170
<211> 114
<212> DNA
<213> Homo sapiens

<400> 170
agcagtggtgt cctccatgaa taaacaggag ttctggaggc ccatcttctg catcttctgc 60
tgattttct tccccatatt tacttaaattc ccacacattc aggccggcggt cagt 114

<210> 171
<211> 150
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(150)
<223> n=A,T,C or G

<400> 171
actgagagca tttataatct gaccaaattc ataggcatta ttaggcttgg ctatcggaag 60
tttctcaggg tcttctggng acctgctgct tttgcctccc ttctcanaag caaggcatcc 120
catggagacc tcccctgcag ggcttccagg 150

<210> 172
<211> 435
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(435)
<223> n=A,T,C or G

<400> 172
atttgttttc cactgcctca cactagttag ctgtgccaaag tagtagtgtg acacctgtgt 60
tgtcatttcc cacatcacgt aagagcttcc aaggaaagcc aaatcccaga tgagtctcag 120
agagggatca atatgtccat gattatcttcc tggtttaggt ctacagtcaa tgtgtatggtg 180
gtctttgtttt cccagtctgc cagaatatct ttgtgcattt ctaatcatttgc gctttaaagc 240
taatcaatgt gttggcagca tctctgtcac tcttgcatttac cacgtgaaga aatcaggttag 300
atttttttct gtggcattgt tttcgaccc aaaatcaggt atgctgacta tttccaaagg 360
gtttttcagt tgcttcattt gcttgtaaag caggaaatcc tcttgcatttgc tttcttttc 420
tcgatgagcc cgtgt 435

<210> 173

<211> 622
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(622)
<223> n=A,T,C or G

<400> 173
actgnnttcc cccaagtcca tgacatgtat acataattaa tggttgcct ccttgattgt 60
tttctccaac atccagacat agaggctgac caacgccttt aatgttatcca gatataacag 120
gattaaggc tcggcacatac acctctggat aaatgttggt cagataccat gtaaaaatttt 180
tacactgaag gcgggtgttt attcaaatac ttttggaaag atcaccaaat gcttttgg 240
taacaatttt tgctgcatac gtatttctcc tataaaaat ttccttgat tcatccatcc 300
agacttctgc aaggcgaact tggttcttag caatcacctg agtgccttt ggaaagctat 360
gagggctttt gctgcgaaaa acatgtccaa caacagagca aggcatatac tccaactgcc 420
caccacattt ccataactctg aaagacattt ctatatttc acctccccag atttccattt 480
cttcatacata gtttccaaata tactcaaaaat attctttga tatggaaaaa agtccctcctg 540
caaaaagtggg tgtttaatt gggtaggggtt catcttcct tctttgcttc tcatacgatcag 600
gaagcgactt ccacccaaatg aa 622

<210> 174
<211> 362
<212> DNA
<213> Homo sapiens

<400> 174
acgggtgcagt tgacccactg ttggctctcc ttgcagttcc tgatatgtca tcttagcat 60
gtggctactt acgtaatctt acctggacac tttctaattt ttggccgcaac aagaatcctg 120
caccggat agatgctgtt gagcagattt ttccctacctt agttcagctc ctgcatacatg 180
atgatccaga agtgttagca gatacctgct gggctatttc ctaccttact gatggtccaa 240
atgaacgaat tggcatggtg gtggaaacag gagttgtgcc ccaacttg 300
gagttctga attgccaattt gtgactcctg ccctaagagc catagggaaat attgtcactg 360
gt 362

<210> 175
<211> 486
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(486)
<223> n=A,T,C or G

<400> 175
acagntnctc tactacactc agcctttat gtgccaagtt ttttttaag caatgagaaa 60
ttgctcatgt ttttcatctt ctcataatcat cagaggccga agaaaaaacac tttggctgtg 120
tctaaaactt gacacagtca atagaatgaa gaaaattaga gtagttatgt gattattca 180
gctcttgacc tggccctt ggtgcctt ggtctgaat ctcccaaaaga gagaaaccaa 240
tttcttaagag gactggattt cagaagactc ggggacaaca tttgatccaa gatcttaaat 300
gttatattga taaccatgtt cagaatgag ctattagatt cattttggaa aatctccata 360
attcaattt gtaaactttt gtaagacctg tctacattgt tatatgttg tgacttgagt 420
aatgttatca acgttttgtt aaatatttac tatgttttc tattagctaa, attccaacaa 480
ttttgt 486

<210> 176
<211> 461
<212> DNA
<213> Homo sapiens

<400> 176
accctggcca ctcccttcct tttggctggc caatgtctcc tctgttaggct ccagaaggct 60
ctcaggatg caggcgccct cctgcagggt tgagttgcaa tggaaacaaa gacagctgtg 120
gtccccatagc accctcatct ggtgacatcc tgctactgac agtcaaaaaga agccttccca 180
gatgaaattt tagtcctctg cgccagccatg ctcttcttcc agcaaaaagag ccatgtgcag 240
tcgggtctgc tccccatggg ggcttgcatg tggggccagc agtggatcag cttccagac 300
acgctcaact ctgcacactc ttccctgccgc ctcaggctt ccaggaccct cccgagcctt 360
atcagagtcc ttacccttag ggctactgat accttgcgtgg gtgaccttg acagattcac 420
ttacctggac tcagttcat aatatgaaaa tgatagggtt g 461

<210> 177
<211> 234
<212> DNA
<213> Homo sapiens

<400> 177
acacatttttaatttacctt ttttgttgtt ttgttagcaac catttggaaa acattccaaa 60
taattccaca gtcctgaagc agcaatcgaa tccctttctc actttggaa ggtgactttt 120
caccttaatg catattcccc tctccataga ggagagggaaa aggtgttaggc ctgccttacc 180
gagagccaaa cagagcccag ggagactccg ctgtggggaaa cctcattgtt ctgt 234

<210> 178
<211> 657
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(657)
<223> n=A,T,C or G

<400> 178
gagctcggan ccctagtaac ggccgccagg gtgctgnat gngcccttc gagcgnncg 60
cccgccagg nacttnatc cccccatc ttccctgtac tcatttgnt ctctcatttt 120
ttggcatatt ttcaagtca cactaaaaa ctcttccatg tattcacttc tcattactt 180
gtctacatgc cgaacctaaag gtcaggattc caaaaagatg agtatactt caaacgcctc 240
ctaaggctt ggtatacatg actttggctg tgcacttcat ttagacttca ctttttgg 300
tgctgttgg ttttacacta gattcccttgc ttccatcaa agataatgaa agattcacat 360
cacagtgcag ctcttcgctt tgcctttcg taagtccgtt gcaactgccg agagttctgg 420
tctgttaggc atgtgtgaaa tccgctttgt ggctctctgt gatttggcc gcttaacgtt 480
tttatttgc ttatttacac atgccaaggt ggcaacgtga aaaatgtctc tgacgctatt 540
ttccgactgt aaagctgagc attcgatata agtagctgct ccaatctgtt tggccatact 600
tgccccctgg tcataggaca ctggcgtctg cctgtgattt gagagctcta ctaatgt 657

<210> 179
<211> 182
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(182)
<223> n=A,T,C or G

<400> 179
acaaaanctt ttaaattttt tattatgg aaactttgct ttgggtttgt ggcaccctgg 60
ccaccccatc tgctgtgac agcctctgca gtccgtggc tggcagttt ttgatcttt 120
aagttcctt ccctacccag tccccattt ctggtaaggt ttctaggagg tctgttaggt 180
gt 182

<210> 180
<211> 525
<212> DNA
<213> Homo sapiens

<400> 180
acacgctttt ggccccgacc aatgaggcct tcgagaagat ccctagttag actttgaacc 60
gtatcctggg cgacccagaa gccctgagag acctgctgaa caaccacatc ttgaagtcag 120
ctatgtgtgc tgaagccatc gttgcggggc tgtctgtaga gaccctggag ggcatgacac 180
tggaggtggg ctgcagcggg gacatgctca ctatcaacgg gaaggcgatc atctccaata 240
aagacatcct agccaccaac ggggtgatcc actacattga tgagctactc atcccagact 300
cagccaagac actatttcaa ttggctgcag agtctgatgt gtccacagcc attgacctt 360
tcagacaagc cggcctcgcc aatcatctc ctggaagtga gcggttgacc ctcctggctc 420
ccctgaattc tgattcaaa gatggaaccc ctccaatgt tgcccataca aggaatttgc 480
ttcggaaacca cataattaaa gaccagctgg cctctaagta tctgt 525

<210> 181
<211> 444
<212> DNA
<213> Homo sapiens

<400> 181
acaccacaat gtgcataaag gagacgtgcc gattgattcc tgcagtcgg tccatttcca 60
gagatcttag caagccactt accttcccag atggatgcac attgcctgca gggatcacgg 120
tggttcttag tatttggggt cttcaccaca atcctgctgt ctgaaaaaac ccaaagggtct 180
ctgaccctt gaggttctct caggagaatt ctgatcagag acaccctat gcctacttac 240
cattctcagc tggatcaagg aactgcattt ggcaggagtt tgccatgatt gagttaaagg 300
taaccatttc cttgattctc ctccacttca gagtgactcc agacccccc accgccttta 360
ctttccccaa ccattttatc ctcaagccca agaatggat gtattgcac ctgaagaaac 420
tctctgaatg tttagatctca gggt 444

<210> 182
<211> 441
<212> DNA
<213> Homo sapiens

<400> 182
acaaccttta ttgcttctcc agcattttcc agaagaatgg tgcatttgc gggccacagg 60
ggatggggga gtaaaaaata acataaacga actgaacaga aatgcaggag ggtggcaaga 120
ggggccgaga ttgggtttc agggcagaga ggtggaaagac cagggcagt cagtgccttct 180
tagctttcag ccaccagat ggagaattcg tcaacccaa ttttgcgtc cccatctttg 240
tctccagcag ccatcagcat cttgggttct ttagcagaca ggtctctggc atctggggag 300
aagccttta ggtatgaatcc cagctcatcc tcctcgatga agccactttg tcctgtcca 360
gcatgtgaaa caccttcttc acatcatccg cactctttt cttcaggccg accatttgaa 420
agaactttt gtggtcgaag g 441

<210> 183
<211> 339
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(339)
<223> n=A,T,C or G

<400> 183
tgtntcatcn taaggggatt gggctctaga tctgtcgacg gcgcatttag gatttgcnat 60
cggttangtg gtccgcgagt catgaatttt tgctctggag cgttattgtt tgtgaagttt 120
atccaggaga gaactatgat tgtgtcgatg cgtttactgc aggaagantc acggctctcag 180
tcacggaggt gtaagggtgg actgactgan tgagacaagg gatatntngt tnttatannc 240
ttgtgatgaa cctgcctacc gtttatgtct ctggctcaat gggtctcng tnctgtnatt 300
cncncaagct gcgggggctt ccnccgttctt gggctctga 339

<210> 184
<211> 490
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(490)
<223> n=A,T,C or G

<400> 184
atatacgcaag cttgtacgac cgacacatac ggccgattgt gctggattgc ttatcttgc 60
gcccgcacgtc tatataancg anactacata gtctcgaaaa tccactcant ttcaagttcc 120
caaaaanacng gaaaaaaacc catgccttat ttaactaanc atcagctcgc ttctccttct 180
gtaccgcgc ttntngctcc cagcctatag aagggtaaaa cccacactcg tgcnncagtc 240
atcnnataac tgattcgccc gggtaactgcc gggcggcgct cganaccaat tngcanaatt 300
cacacattgc ggcgctcnan aagctctaga aggccaatcg ccataattgat ctatacatta 360
tggccgtcgt tnacacgtcg tgacgggana ncctggngta ccattaatcg ctgcacantc 420
ccttcgcagc tgggtntac aaaagccgccc catcnctcca cgttgcncc gatggcaagg 480
acnccctnat 490

<210> 185
<211> 368
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(368)
<223> n=A,T,C or G

<400> 185
ctnnanatag cangcttcta cgaccgacac aatacggcca ntgtgcttggaa ttgccttcag 60
cgccgcggcgg gcagtacgg cgctcatcta tcngatgatg ggcaccaat gtgggtttt 120
aacctttta tatggctggg gacanaaagc gcggttacnn aaccnataac gagctgatgg 180
tcatttaaaa atgcttgggg tttcccggt ctttgggaa attgaaactg agtggactt 240

canaaaactgt gctactttcg cttatctaag tactcggccg caacacccatg ccgaatccgc 300
anatatcatc acnctggcg gcgtcancat gcntctaaag ggccaattcn cctanatgag 360
tcttatac 368

<210> 186
<211> 214
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(214)
<223> n=A,T,C or G

<400> 186
ngggagatcg cagcttgta gactcgtcat ataacgnca atgtgctgga tcgcttcanc 60
gccggccggcg gtctaattcg gttcggattt tttgtgtntt gtctntntta canggtgcta 120
tcccccttctt cctccttc tggccatcctc atcctttatc tccttttgg acaagtgtca 180
nancagacag angcagggtg gtggcaccgt tgaa 214

<210> 187
<211> 630
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(630)
<223> n=A,T,C or G

<400> 187
cagctggac gagtcgatca tatacggcgc atgtgttgna tcgctatcgt gtccggcgag 60
tantattan attactgtta ttctctgtcc tactggatat gatcttttga nggcangtct 120
gtgtcgtctg gtcacaccat gttctcaggg tggcaaaata ctttcctata atagtttatg 180
gataatgaat gacgactang tctanaaana cgctagctaa ataacacact cagggaaaga 240
gtcttaaata ttgtgaaggt gtttttanta tacaacnttt gtttacataa tagaaataa 300
tttttagact tttaaacaga cacttgagcc agatttttta atgttaccat ctatagtgtc 360
ttgaaaatat tcctcttagt ttccaatatg aatgaatcta aaatccatct tttcaattat 420
gcccaggccc gtggtcaatg cnccctcnac acttcattaa cggattatac cttggaaac 480
cataatctgg ctnaggacga atcgccctggc ncangctaann aactgccctg tattgagggg 540
ttatnnctga ttgcngaggt gcctctccag gtccccaaag ggtcgtaactg ttgaanctgg 600
ctctaatttt ntcttgccctn acaggtctcc 630

<210> 188
<211> 441
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(441)
<223> n=A,T,C or G

<400> 188
cnngcaanac anggtcggat tccgntgagg naanaattcc ctnatagggc tcgcccccta 60

ttcaccaaac caancngaaa ctcttgcggc caaatctaag ctatnnncaca accccactct 120
gnagggtatg cgccccggcc ctgcaatgaa atcaatanca tatttgaga cagagagata 180
gagagagaga ggttcctggc cttnnctatt ctgctttac ttgnnagatn tcaganatag 240
aaaaacctat ccttaggtccn nccaatgatn gcggcttncg aatcccgng tggccantcc 300
ccggatcgga ctaaatcaa gaagatcctc cgtcntcctg ttcctccaca ctggagtccc 360
attgtatgca tgggtnttc actggctnat cataccnnag gatctgtcca ccttnaactc 420
ttctctngga antccctncc c 441

<210> 189
<211> 637
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(637)
<223> n=A,T,C or G

<400> 189
agggnngtata tacccacttg tacnactcga tcatanacgc gcatntctga atcgcttnct 60
ggccgcgatg tactgtggc acttaagcac tgagtactgt ttgcgtcatg ccngtcan 120
agatgctgct gcaaaggac tccaaacnaaa tacactgtct tcaacaggag ttaacacetc 180
acacttggtg ganaanagaa ctcaactggtg gtgatgcaca cgactgnatc catcaagtgc 240
gtttgcctgt tgactgctaa ccaaggctct ggcagtagct gcccggggcgg cgctcgaaac 300
caaatctgca aatatcatca cactggcggn cgctcagcat catctanaag gccatcgct 360
atagttagtc tatacatcat ggcgcnttt acactcctac tggaaaacct gcgtaccact 420
taatcgcttc acacatcccc ttgcngtn gcttatancn aaaagccac gatgcctcca 480
cattgcncnc tgatggcatg ancccctac ggcataanc gcggtntgt tacncangt 540
accgtntgc acgctacnch tttccttct cctttcccc ttcccgttcc tcaccattcg 600
gggccttagg tcnatatctc gnccacccaa atntagg 637

<210> 190
<211> 653
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(653)
<223> n=A,T,C or G

<400> 190
agggggtata tacccacttg tacgactgna tcataatacgc gcatgtctgg aatcgcttnc 60
gtggctgcca tgtattgaca ctacttctaa gaactacaaa agtgatactg angatacatt 120
acacagaang gctnacatc tcncagatcc tcattttca tggatgtgg acatcangan 180
cacgtggata agtgtatcta aanaatggct ttcaaaaat tttcaacttta ttaaggttt 240
acatganatt cataaaatgt cttaatacta tttctnaaaa taacatctaa tcggaaaacta 300
tgcctnaact gcacntttt tgtgtanata atcntanttg tacgcccggc ggcgcacaaag 360
ccnaatctgc gattcctcac ctggcgccgc tcaacatcat ctaaaggcca atcgctata 420
ntantctata catcctggcc gcgtttacac gtctaattgg aaaccggcgt accacttac 480
gcgtgcagca ctcccccttcc cactgggtta tacnaaagcc gcgcgtatgcc tcccacattc 540
canctgatgc aatgaccct gttcgcctta ncccgccgtt tggatccca ntnaccacnt 600
cagcgctgcn cntcttctt ctccttcttgc ccnttncgt tccctcactc nng 653

<210> 191

<211> 663
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1) ... (663)
<223> n=A,T,C or G

<400> 191
anggngtata tacccactgt ncgactcgat catatacgcg catgtcggat cggctccanc 60
gcccggcat gtactatatac tacatcaact gtattatcat ttanatattt atnaaaagaca 120
aaatcatact tccatctgct cactgatgat aattactatg atacatgatc atgtaaacgt 180
atcaatataa caatggaaga tccctctgac tatgcaagcc taatttcca atcnatgca 240
ctctcatagc tcaaananatn cacngacatc ctgatgaaac tatnatacan tttccacaca 300
aatcacttcg ctttagatct ctccattatt cttgctttc ccccctaaca actacaaatc 360
ctcntggat gggagaata tatacatct actaaaaata atatataatc ccctgcanat 420
ttgtggnaaa tcnggtgtct caanagccac aggagnacaa ggggnacca actaggactt 480
ttgtatgctt atctctgtac tcgacacac ctaagcgatt ctgcnattct ccctggcggc 540
gtcacanctc tanaggccat cncnatatga tctatacatc ntggcgtctt tacactctga 600
cgaaaaccgg gtnccantta ccctggacca tcccttcgcn ctgntataca aagcccccg 660
ncc 663

<210> 192
<211> 361
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1) ... (361)
<223> n=A,T,C or G

<400> 192
anttttata tacccactgg tacaactcgaa ncctatacgg cgcantncg gaatcanctt 60
cancggcgcc ggcattgtacc ggtnatcatc atcngatgat ggcgctcnaa tgtgggtttt 120
acctnttata cggctgagat canatcgcgt acataacaaa nncaactgat ggttaatnta 180
aatncggtt ggttctcccn ntctgttggg gaacttgana ctgagtgnga cntccatana 240
cgtgctattt tcggctancn antcctcagc gnacacccat ngnagtgcgc naattcatcc 300
atgntggcct cgactnttcc aaaangccnt ncgcccacnt gntcgcnana cantctcggc 360
c 361

<210> 193
<211> 314
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1) ... (314)
<223> n=A,T,C or G

<400> 193
agggngnata taccaactgg tncgactcgaa tcctatacgc gcatttcgga ttgcgttcaa 60
cggcgccggc atgtaccaaa cctcaatccc aaccgtctca ntngacggg ctcaggctt 120
tcacagccac cccacatttc tttgttttgc tctgccactt caaaagaatt ccaaataaga 180

attctgctgc agctccgtac aaggatatgg gcagcacagc acacacagag tngtgctcct 240
cacacttctc tggnaatgtc tcgtgaatat ctcaacagtc angaagtggg gcgttatcaa 300
aaacaatcag ggcc 314

<210> 194
<211> 550
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(550)
<223> n=A,T,C or G

<400> 194
aggngngata tacccactgg tncgactcga tcctatacgc gcatgtcgga ncgctatgtg 60
gtcnccgaag tacctcttgc gcaatgtatgg tctgtntcct ctatgatnag tgatcgaata 120
atcatcgaat tcancgaaag ttatcgagt gatatntgtg gctttagaa tctatgctcc 180
atggtgtggt cactgtcaag attaacacag aatggaaagan ncngcactgc ataaaagatg 240
ttgtcaaatt ggggcgttg atcngatagc tcntcccaag aggtcantgg tgttcaggat 300
tnccnacataa gatnttgat cacngacga ccagangata ccngtgcaa ctgtgaancn 360
ngtaatctgc ctatnccgc cctctggan gatccctcg ggacgacgag atcattctgg 420
aaacagcnn tgatagtcca gtnnangatt gatgancgac ganacgcntg atanatgtct 480
gacgtgagat tnggatgtga atcttccnt gtgtgacctg cnccntaccn aanggtgcgn 540
ctccactcnn 550

<210> 195
<211> 452
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(452)
<223> n=A,T,C or G

<400> 195
nngcgggnat gataccaact ggtacgaact cganctctat nacggcgctn tttcnngatc 60
tgctatgtgg tctcggaat gtacattata acngggcana catataatct acntctgtct 120
ttntctcccc cngagagcgc aancatctcc aaatcgggtt ctgggtcatc caatggtctc 180
cantaatcac acaactcata tatattttagt gaangtgtct gtcatcgtcc ccacgangga 240
agtnncgtcg ctgtntgtct gtcacttaggt gngtactctc cagtaattga aanctggtna 300
nggctgtctg tngtactggc cggccccc gaaancaat ctgtnnatcatcacatng 360
cgncccccga ncatcactna gggncanttc gcctatactg atcgtntgcg annctgcgn 420
cncttacacg tcgnacggga naccggcctt cc 452

<210> 196
<211> 429
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(429)
<223> n=A,T,C or G

<400> 196
gcgggnnnat gataccagct ngtacgactc gatccataa cggcgcatgt gngtatccgc 60
tacgtgtctc ggcgatgtac atataacggg gcaacatata atnatacant ctgtcttttt 120
ctccccccga aacggcaacc atctccaata tcggctctggg tctccaatgg tctccaacta 180
aatcacacaa gtcaaataata ntangaaa gtgtctgtct cntccccaga aggagtancg 240
ttagctttg tctgtcatta gggtggtacc tccagtnaca tgaaaactgg tgagggtgtc 300
cttgtacaag ctctgcctca ccagatccta tactattagg gggcccacgg ttatctatct 360
taaggtctn aaaacctgga cttcatctgc tccggcggan gaatgtcccg cttacttacg 420
ntgtccac 429

<210> 197
<211> 471
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(471)
<223> n=A,T,C or G

<400> 197
atgatacgcgac gctngtacga gccgtcaacta tnacggcnca ttgtgtggat tcngctntga 60
tcggcgcccg ggcatgtcca tcnagagcgc atcatgggan tgnactcccc atatnntgac 120
caangttcgc gcaaggagcc nagancggat actacctgag ctgtcgctn gttatacacg 180
tttctggcca angancaact ccacatncaa caagttggtg ttgaaatgtt gtttatnagt 240
ccaccaaccg gccgctctgt cccttcccga tgatccgaag ataagcttcc tgccggaaan 300
acgaacggcg tgggtgtgngg acatantgat atgtcggtt caggaagtac tcgnccgcaac 360
ncgcaagcna atctgcnata tcacacactg gccggcgtcg agctgccana ngcccnntcg 420
cctatatgat tctatacatt cctggccgtc tnttacactc ngacgggaaa c 471

<210> 198
<211> 643
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(643)
<223> n=A,T,C or G

<400> 198
tngtncgacc gtcactatac gcccattgtgt ggatccgntc cacggcgccg ggcangtacg 60
anactatatt gatcctctga tattgaaagt tggctcana ataacctta angcaaata 120
ctcantgagt tttgaccaga agtcaccaca tcatgaatca cagtctatgg caaatgatac 180
cagttctct aagtccatg ctcaaggtaa gagcatgcta ttccgtttta catttactgg 240
aatttactgt tcattcatna ttaaaatctc tagtttcat cctcaactgt ctaanaccag 300
tgtgcacaga cttaagactc tggctcctc attttctcca acagaaacat tctcagtgtc 360
tactgttcta aaaggaaatt tccgaggtgg cacttctcg aatatcgacc ctenggctct 420
atcaggcggtt acttcnnnca ctcgtcattt gggctgttc anttgcattt tctgtccagt 480
cacttcattt taagaaaaca attgatcgct ggtcacatgt nattcattgg cagccgggtgt 540
gactgctgag tctcgccgac acncttagcaa tcgnnattct ccatggngcg tcactctcta 600
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<210> 199

<211> 292
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(292)
<223> n=A,T,C or G

<400> 199
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gtccggcgag tctatgtat ttatttnnga ttaaatcaat attttcttc tgaatattaa 120
tcttatctnt acttttatac tattgaccta gctatatgta ttgancttt tgaactccta 180
tcagtnttt tcatactatc gtatatttc cacttggta ctntngctga ntccctagata 240
tcgtaaaaca tctctnnatc ntcacacnnga gnccagggn tctgtatngaa tt 292

<210> 200
<211> 275
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(275)
<223> n=A,T,C or G

<400> 200
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tgcttancgt ggtcnccggcc gaagtactat gctatnttac tttttggga tataaaatca 120
atataattct ttctnaagta tataaatctt atccnctatc cnntcnatac ctntctgaca 180
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tganntcttc cacnttggta cctttacgc tgaat 275

<210> 201
<211> 284
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(284)
<223> n=A,T,C or G

<400> 201
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atctatctcg actgattcac ctgtcattgt aaanaattcg tgcagctgt ctaccnctta 120
nacatcatct aatcnaacta ncgtataaa tttcttcaat agggatanac ntntagtaca 180
tacgnttcca ttgagntacn tccgcggacc cncatcgcaa acnncatgcg gtcagtcnna 240
gcacccctca tcttaatccg tccttacnt ntgaacgctc cact 284

<210> 202
<211> 448
<212> DNA
<213> Homo sapiens

<220>

<221> misc_feature
<222> (1)...(448)
<223> n=A,T,C or G

<400> 202
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ttcgacggac gccgggcatg tactttata atnctactcc tcagaccttg catctcnacc 120
gctnggtcca gttttaaaa acnnacttcc gtngtgcagc cctggttctg ancantctct 180
atcacnctct atcctcnat ccncaanact anatcgctg aattcatatt tattcatttt 240
ccataatgat ggggaaanga ctatcnatna tnatgcttan cacnctngct gcanttcgnc 300
natctcgcnna ncntgaaac gattactctg tcgcgaaccc tctangntga attctgcnaa 360
atatctntna cnctggcngg cgctcnangn atgcctctcg anggccaatc cgccnngcat 420
gattctaatt anatccntng gtcccnntt 448

<210> 203
<211> 321
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(321)
<223> n=A,T,C or G

<400> 203
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tgtccggcga ngtaccatataatcgaanta ncataggttct ggangcccnc tcattttcaa 120
tttcccaaaa nacgggaaaa ccnaagcctt atttaactaa ctatctgctc gcttctcgct 180
tctgtaccgc gctatntgtt nccagcctat aanaaggta aaacccacac tcggtgcgtc 240
agtctccnat atantgagtc nccgggtact ggccgggcgg tcgttcnaaa ncaattcneg 300
aanttcacta ctggcggcgc c 321

<210> 204
<211> 369
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(369)
<223> n=A,T,C or G

<400> 204
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acttgcgcg gccaagtatc tataaagcaa actatcacag ttctgaaagt ccatctcant 120
ttcagttccc aaaagancgg gaaaaacccaa gccttattaa actaacaatc agtcgctctc 180
gcttctgtac cgcgttttg gccccccagcc tataaaaggtaaaaacccac actcggtgcg 240
ccagtcatcg ataactgaat cgcccggtac tgcccgccg ggcgctcnann ccaaattctgc 300
agatatcaca cactggcggc gctcancatg ctctagaagg ccaattcncc tatantgatt 360
ctattacaa 369

<210> 205
 <211> 2996
 <212> DNA
 <213> Homo sapien

<400> 205

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cctacaccc	ggacaggggac	agtctctatg	tcaatggttt	cacacaggcg	agctctgtgc	180
ccaccactag	cattcctggg	accccccacag	tggacctggg	aacatctggg	actccagttt	240
ctaaacctgg	tccctcggct	gccagccctc	tcctggtgct	attcactctc	aacttcacca	300
tcaccaacct	gcggtatgag	gagaacatgc	agcacccctgg	ctccaggaaag	ttcaacacca	360
cggagagggt	ccttcagggc	ctggccctg	ttcaagagca	ccagtgttgg	ccctctgtac	420
tctggctgca	gactgactt	gctcaggct	gaaaaggatg	ggacagccac	tggagtggat	480
gccccatctgca	cccaccaccc	tgaccccaaa	agccctaggc	tggacagaga	gcagctgtat	540
tgggagctga	gccagctgac	ccacaatac	actgagctgg	gcccctatgc	cctggacaaac	600
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gatctgcaat	gactggaact	tgccggtgcc	tgggtgcct	ttcccccagc	cagggtccaa	2940
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<210> 206

<211> 914

<212> PRT

<213> Homo sapien

<400> 206
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Asn Leu Val Pro Arg Leu Pro Ala Leu Ser Trp Cys Tyr Ser Leu Ser
35 40 45
Thr Ser Pro Ser Pro Thr Cys Gly Met Arg Arg Thr Cys Ser Thr Leu
50 55 60
Ala Pro Gly Ser Ser Thr Pro Arg Arg Gly Ser Phe Arg Ala Trp Ser
65 70 75 80
Leu Phe Lys Ser Thr Ser Val Gly Pro Leu Tyr Ser Gly Cys Arg Leu
85 90 95
Thr Leu Leu Arg Pro Glu Lys Asp Gly Thr Ala Thr Gly Val Asp Ala
100 105 110
Ile Cys Thr His His Pro Asp Pro Lys Ser Pro Arg Leu Asp Arg Glu
115 120 125
Gln Leu Tyr Trp Glu Leu Ser Gln Leu Thr His Asn Ile Thr Glu Leu
130 135 140
Gly Pro Tyr Ala Leu Asp Asn Asp Ser Leu Phe Val Asn Gly Phe Thr
145 150 155 160
His Arg Ser Ser Val Ser Thr Thr Ser Thr Pro Gly Thr Pro Thr Val
165 170 175
Tyr Leu Gly Ala Ser Lys Thr Pro Ala Ser Ile Phe Gly Pro Ser Ala
180 185 190
Ala Ser His Leu Leu Ile Leu Phe Thr Leu Asn Phe Thr Ile Thr Asn
195 200 205
Leu Arg Tyr Glu Glu Asn Met Trp Pro Gly Ser Arg Lys Phe Asn Thr
210 215 220
Thr Glu Arg Val Leu Gln Gly Leu Leu Arg Pro Leu Phe Lys Asn Thr
225 230 235 240
Ser Val Gly Pro Leu Tyr Ser Gly Cys Arg Leu Thr Leu Leu Arg Pro
245 250 255
Glu Lys Asp Gly Glu Ala Thr Gly Val Asp Ala Ile Cys Thr His Arg
260 265 270
Pro Asp Pro Thr Gly Pro Gly Leu Asp Arg Glu Gln Leu Tyr Leu Glu
275 280 285
Leu Ser Gln Leu Thr His Ser Ile Thr Glu Leu Gly Pro Tyr Thr Leu
290 295 300
Asp Arg Asp Ser Leu Tyr Val Asn Gly Phe Thr His Arg Ser Ser Val
305 310 315 320
Pro Thr Thr Ser Thr Gly Val Val Ser Glu Glu Pro Phe Thr Leu Asn
325 330 335
Phe Thr Ile Asn Asn Leu Arg Tyr Met Ala Asp Met Gly Gln Pro Gly
340 345 350
Ser Leu Lys Phe Asn Ile Thr Asp Asn Val Met Lys His Leu Leu Ser
355 360 365
Pro Leu Phe Gln Arg Ser Ser Leu Gly Ala Arg Tyr Thr Gly Cys Arg
370 375 380
Val Ile Ala Leu Arg Ser Val Lys Asn Gly Ala Glu Thr Arg Val Asp
385 390 395 400
Leu Leu Cys Thr Tyr Leu Gln Pro Leu Ser Gly Pro Gly Leu Pro Ile
405 410 415
Lys Gln Val Phe His Glu Leu Ser Gln Gln Thr His Gly Ile Thr Arg
420 425 430
Leu Gly Pro Tyr Ser Leu Asp Lys Asp Ser Leu Tyr Leu Asn Gly Tyr
435 440 445
Asn Glu Pro Gly Pro Asp Glu Pro Pro Thr Thr Pro Lys Pro Ala Thr

450	455	460
Thr Phe Leu Pro Pro Leu Ser Glu Ala Thr	Thr Ala Met Gly Tyr His	
465 470 475	480	
Leu Lys Thr Leu Thr Leu Asn Phe Thr Ile	Ser Asn Leu Gln Tyr Ser	
485	490	495
Pro Asp Met Gly Lys Gly Ser Ala Thr Phe	Asn Ser Thr Glu Gly Val	
500	505	510
Leu Gln His Leu Leu Arg Pro Leu Phe Gln	Lys Ser Ser Met Gly Pro	
515	520	525
Phe Tyr Leu Gly Cys Gln Leu Ile Ser Leu	Arg Pro Glu Lys Asp Gly	
530 535	540	
Ala Ala Thr Gly Val Asp Thr Thr Cys Thr	Tyr His Pro Asp Pro Val	
545 550	555	560
Gly Pro Gly Leu Asp Ile Gln Gln Leu	Tyr Trp Glu Leu Ser Gln Leu	
565	570	575
Thr His Gly Val Thr Gln Leu Gly Phe	Tyr Val Leu Asp Arg Asp Ser	
580	585	590
Leu Phe Ile Asn Gly Tyr Ala Pro Gln Asn	Leu Ser Ile Arg Gly Glu	
595	600	605
Tyr Gln Ile Asn Phe His Ile Val Asn Trp	Asn Leu Ser Asn Pro Asp	
610 615	620	
Pro Thr Ser Ser Glu Tyr Ile Thr Leu	Leu Arg Asp Ile Gln Asp Lys	
625 630	635	640
Val Thr Thr Leu Tyr Lys Gly Ser Gln	Leu His Asp Thr Phe Arg Phe	
645	650	655
Cys Leu Val Thr Asn Leu Thr Met Asp	Ser Val Leu Val Thr Val Lys	
660	665	670
Ala Leu Phe Ser Ser Asn Leu Asp Pro	Ser Leu Val Glu Gln Val Phe	
675	680	685
Leu Asp Lys Thr Leu Asn Ala Ser Phe	His Trp Leu Gly Ser Thr Tyr	
690 695	700	
Gln Leu Val Asp Ile His Val Thr Glu	Met Glu Ser Ser Val Tyr Gln	
705 710	715	720
Pro Thr Ser Ser Ser Thr Gln His Phe	Tyr Leu Asn Phe Thr Ile	
725	730	735
Thr Asn Leu Pro Tyr Ser Gln Asp Lys	Ala Gln Pro Gly Thr Thr Asn	
740	745	750
Tyr Gln Arg Asn Lys Arg Asn Ile	Glu Asp Ala Leu Asn Gln Leu Phe	
755	760	765
Arg Asn Ser Ser Ile Lys Ser Tyr Phe	Ser Asp Cys Gln Val Ser Thr	
770 775	780	
Phe Arg Ser Val Pro Asn Arg His His	Thr Gly Val Asp Ser Leu Cys	
785 790	795	800
Asn Phe Ser Pro Leu Ala Arg Arg Val	Asp Arg Val Ala Ile Tyr Glu	
805	810	815
Glu Phe Leu Arg Met Thr Arg Asn Gly	Thr Gln Leu Gln Asn Phe Thr	
820	825	830
Leu Asp Arg Ser Ser Val Leu Val Asp	Gly Tyr Phe Pro Asn Arg Asn	
835	840	845
Glu Pro Leu Thr Gly Asn Ser Asp Leu	Pro Phe Trp Ala Val Ile Leu	
850	855	860
Ile Gly Leu Ala Gly Leu Leu Gly Leu	Ile Thr Cys Leu Ile Cys Gly	
865 870	875	880
Val Leu Val Thr Thr Arg Arg Arg Lys	Lys Glu Gly Glu Tyr Asn Val	
885	890	895
Gln Gln Gln Cys Pro Gly Tyr Tyr Gln	Ser His Leu Asp Leu Glu Asp	
900	905	910
Leu Gln		

<210> 207
<211> 2627
<212> DNA
<213> *Homo sapiens*

<400> 207

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tagcatcatc attattctgg ctggagcaat tgcaactcatc attggctttg gtatttcagg 180
gagacactcc atcacagtca ctactgtcgc ctcagctggg aacattgggg aggatggat 240
cctgagctgc acctttgaac ctgacatcaa actttctgtat atcgtgatac aatggctgaa 300
ggaagggttt ttaggcttg tccatgagtt caaagaaggc aaagatgagc tgccggagca 360
ggatgaaatg ttccagaggcc ggacagcgt gtttgcgtat caagtgatacg ttggcaatgc 420
ctctttcggt ctgaaaaacg tgcaactcac agatgctggc acctacaaaat gttatatcat 480
caactctaaa ggcaggggaa atgctaacct tgagtataaa actggagcc tcagcatgcc 540
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tcaagagaat gattaaatatac acatttcata caccaaaaaaaaaaaaaa 2627

2627

<210> 208
<211> 282
<212> PRT
<213> *Homo sapiens*

<400> 208
Met Ala Ser Leu Gly Gln Ile Leu Phe Trp Ser Ile Ile Ser Ile Ile
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Ile Ile Leu Ala Gly Ala Ile Ala Leu Ile Ile Gly Phe Gly Ile Ser
20 25 30
Gly Arg His Ser Ile Thr Val Thr Thr Val Ala Ser Ala Gly Asn Ile
35 40 45
Gly Glu Asp Gly Ile Leu Ser Cys Thr Phe Glu Pro Asp Ile Lys Leu
50 55 60
Ser Asp Ile Val Ile Gln Trp Leu Lys Glu Gly Val Leu Gly Leu Val
65 70 75 80
His Glu Phe Lys Glu Gly Lys Asp Glu Leu Ser Glu Gln Asp Glu Met
85 90 95
Phe Arg Gly Arg Thr Ala Val Phe Ala Asp Gln Val Ile Val Gly Asn
100 105 110
Ala Ser Leu Arg Leu Lys Asn Val Gln Leu Thr Asp Ala Gly Thr Tyr
115 120 125
Lys Cys Tyr Ile Ile Thr Ser Lys Gly Lys Gly Asn Ala Asn Leu Glu
130 135 140
Tyr Lys Thr Gly Ala Phe Ser Met Pro Glu Val Asn Val Asp Tyr Asn
145 150 155 160
Ala Ser Ser Glu Thr Leu Arg Cys Glu Ala Pro Arg Trp Phe Pro Gln
165 170 175
Pro Thr Val Val Trp Ala Ser Gln Val Asp Gln Gly Ala Asn Phe Ser
180 185 190
Glu Val Ser Asn Thr Ser Phe Glu Leu Asn Ser Glu Asn Val Thr Met
195 200 205
Lys Val Val Ser Val Leu Tyr Asn Val Thr Ile Asn Asn Thr Tyr Ser
210 215 220
Cys Met Ile Glu Asn Asp Ile Ala Lys Ala Thr Gly Asp Ile Lys Val
225 230 235 240
Thr Glu Ser Glu Ile Lys Arg Arg Ser His Leu Gln Leu Leu Asn Ser
245 250 255
Lys Ala Ser Leu Cys Val Ser Ser Phe Phe Ala Ile Ser Trp Ala Leu
260 265 270
Leu Pro Leu Ser Pro Tyr Leu Met Leu Lys
275 280

*
<210> 209

<211> 309

<212> PRT

<213> Homo sapiens

<400> 209

His Ala Ser Ala His Ala Ser Gly Arg Gln Arg Gln Leu His Ser Ala
5 10 15

Ser Thr Gln Ile Arg Trp Glu Pro Ser Pro Ala Met Ala Ser Leu Gly
20 25 30

Gln Ile Leu Phe Trp Ser Ile Ile Ser Ile Ile Ile Leu Ala Gly
35 40 45

Ala Ile Ala Leu Ile Ile Gly Phe Gly Ile Ser Gly Arg His Ser Ile
50 55 60

Thr Val Thr Val Ala Ser Ala Gly Asn Ile Gly Glu Asp Gly Ile
65 70 75 80

Leu Ser Cys Thr Phe Glu Pro Asp Ile Lys Leu Ser Asp Ile Val Ile
85 90 95

Gln Trp Leu Lys Glu Gly Val Leu Gly Leu Val His Glu Phe Lys Glu
100 105 110

Gly Lys Asp Glu Leu Ser Glu Gln Asp Glu Met Phe Arg Gly Arg Thr
115 120 125

Ala Val Phe Ala Asp Gln Val Ile Val Gly Asn Ala Ser Leu Arg Leu
130 135 140

Lys Asn Val Gln Leu Thr Asp Ala Gly Thr Tyr Lys Cys Tyr Ile Ile
145 150 155 160

Thr Ser Lys Gly Lys Gly Asn Ala Asn Leu Glu Tyr Lys Thr Gly Ala
165 170 175

Phe Ser Met Pro Glu Val Asn Val Asp Tyr Asn Ala Ser Ser Glu Thr
180 185 190

Leu Arg Cys Glu Ala Pro Arg Trp Phe Pro Gln Pro Thr Val Val Trp
195 200 205

Ala Ser Gln Val Asp Gln Gly Ala Asn Phe Ser Glu Val Ser Asn Thr
210 215 220

Ser Phe Glu Leu Asn Ser Glu Asn Val Thr Met Lys Val Val Ser Val
225 230 235 240

Leu Tyr Asn Val Thr Ile Asn Asn Thr Tyr Ser Cys Met Ile Glu Asn
245 250 255

Asp Ile Ala Lys Ala Thr Gly Asp Ile Lys Val Thr Glu Ser Glu Ile
260 265 270

Lys Arg Arg Ser His Leu Gln Leu Leu Asn Ser Lys Ala Ser Leu Cys
275 280 285

Val Ser Ser Phe Phe Ala Ile Ser Trp Ala Leu Leu Pro Leu Ser Pro
290 295 300

Tyr Leu Met Leu Lys
305

<210> 210
<211> 742
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(742)
<223> n=A,T,C or G

<400> 210
cattgggtac gggccccctc gagtcgacgt atcgataagc ttgatatcga attcggcacg 60
aggcccgacc gtcctcgag agccagcaac gggcagtat gttagccccc gagaaaaat 120
tacatgcgga atggaaagca ggcgctcagg gtggctcctg ctggaatgag agctggagtg 180
caggctccgt gggttcctggg catgcgggtg tggctcagtt ctcaccttgc agatggagtg 240
ggactgttga cccaggccag cctggggact gcctcctcac ctccctgcgc aggctgacct 300
tgtcaccttg cctcttgagc ttgcctctct cctgcccaga ngtccttga gcaaaaatgga 360
ggtcgagagg catttggcac tcacgcctca ccacggacac tggtgcattc ttgggtaccc 420
cttggcctca atctattgct gggggangga ngactgangc ccattgctgg ggcctgaat 480
gcagggactg taaccaccca tccccttctc agggcacctc tccctctcca gcacncttgc 540
tttgcattta atgctaccta atttcctact gangtggctc agaagctctt ccgcatttgc 600
ccttggccgc agcaaatttt tatcccttagg gttaaagataa cagaaggcan ctttgggcct 660
tgcctgccac attctcaggt ntncactgaa gcacagtatc tatttctcca aaaatagggg 720
ctgttaactt gttactaccc cc 742

<210> 211
<211> 946
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(946)
<223> n=A,T,C or G

<400> 211
ggcacgaggc acatcgctgg atttctcatt gccaagctct attaattcat tcttttcat 60
aacctttat tcttatttca tggatgcaac attttcttg tctctcagg aataataatt 120
attcctactt ttaaaggctt aatttctta ttactttatt tctctggag tgagttttc 180
ctaaaggat aatgagatgg aaaatgaaaa aacaaatgg agacatggag ataccttctg 240
aaactcaagc attcctctac gtggatgtgc cagaggaaaa gaacagaaca aaggagggt 300
gacactattt aaataaaaaat atataagaat attacataac aaacaaaaaaa gcccaaattcc 360
tcaggttcaa aaggaggaga aaatgtcaag caagacaaaa acagatgaag caacaaaaaa 420
agtgacatag ctggtcaccc atattgaaat ttcagaacat gagtataaa ggactcccag 480
aaaaaaaaaaa aacccaaact aaaaaacaga aaaaaaggac tttaccaccn aaaacttgan 540
gaatcaggaa gactcagttt ctcattaaga aaantgtat agggatggg ggcaaggct 600
tcaaagtngc aggggatacc aataaacctt ctgaagttt ggaacttcat actccaaaat 660
ngaatttttgc ttgaaatgc cccggtagg ggccaaatttt aggacttaga aaggacccng 720
gnaaatcatt cccnncttgc cccccccgaa agaaattaat agaagggtt tattccgc 780
attannaaaa aaggaatcca ggaattnccg ntttttcca gtgttangnt ggggntgtan 840

aaactgaggg cttagcaagg gcgggnattaa ccacccnngg tcccacccca aaantggnnng 900
gggtgggccca caaattcggg nttntncct ttaangcgaa aaccc 946

<210> 212
<211> 610
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(610)
<223> n=A,T,C or G

<400> 212
ggcacgaggt ttctggctgg agcctcgac actggctcac tgcatgggt ggtgtcgaca 60
gtggtagat ggcaaccagt aacgggagct ttcctgcca ggcaggaga cgagtagaaag 120
ggagcggcat gctggaggct ggagcctgag cccctggggc tcgccttgct gtgttgggt 180
gtgacgtggg acactgcagc tcggccagag tggtaaaaaa tgcctgggt tacgctttc 240
tggcttgcc cgtctatctg ctccaagcca ggctgganga ngagganaag gaatcacctg 300
tggtagctg gagcctgcat gtggcgtgac tctgcaactc gcctcgtgt actgatggca 360
gccacggaga ctgcagctcg acagggagtg aggcttctca ntggcttga agctcagctg 420
actcccacga aatttgcgg aaactcaagg ctgtcagtga ctttcgtggc gccaagactt 480
aancangcgc gttgcgtgca tccggccagt gtctgtgcca cgtgcctgaa cnccacctt 540
anataancac cccgaacgcg cnncgcgcag gccgcgcga cacgnccggg cancaactt 600
gctggcttcc 610

<210> 213
<211> 438
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(438)
<223> n=A,T,C or G

<400> 213
ccganagcgg tttaaacggg ccctctagac tcgagcggcc gccctttttt tttttttt 60
aaataaattt ctagattatt tattacataa gcagaccact gaaacattt ttcaaaagta 120
ttccatggat agtcaaaaac atattgatat gattattatt ggtctgttaa agaaaacaaa 180
ataaaaagaa caaactggga attatcaata aacaaatcaa aacttagatg taattataac 240
ctaaagggtcacagggcaa atgtgaagca agcttctgtc tcagagcctg catatggaa 300
acatgttagta cttagctttg gcatcttct ttcctcctt tggttgagtt taagtattaa 360
taaaaggtgg actgagaaaaa ctttttttta caatcttatg gggatttt agtgaaaacg 420
ttttagaagt aggaatat 438

<210> 214
<211> 906
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(906)
<223> n=A,T,C or G

<400> 214

gccctctaga tcgngcggcc gccctttttt tttttttt gaaataaatt tctagattat 60
 ttattacata agcagaccac taaaacattt attcaaaagt attccattga gagtcaaaaa 120
 catattgata tgattattat tggctgtta aagaaaacaa aataaaaaga acaaactggg 180
 aattatcaat aaacaaatca aaacttagat gtaattataa cctaaagggc tcacagggca 240
 aatgtgaagc aagcttctgt ctcagagcct gcataatggaa gacatgttagt acttagctt 300
 gncatcttc tttcctcctc ttgnttgagt ttagtattaa taaaagttgg actgagaaaa 360
 cctttttta caatctttag ggttattttt agtggaaacg tttagaagta gaatatacat 420
 attaaaactg cncagaacaa atgnggtgca tctcaaattgg nggtccattt tcaaaatatg 480
 aacacatatg ggcagcattt tttttttaa aaagtcagaa ggggcctnct catgcccctt 540
 tccacttctt cactcattgg nccttcaacc caagcttaac tactntcctg acctccaaca 600
 tcataaacta gttccnagc tttgaaacct tttccaatg agtcntaccg gaatagatgn 660
 tcacagaanc ctcttaaaaa ttttggaccc tgcccggnnt ntaaaaaggg tgcaataaac 720
 ccaccaacat ctggctggg ggggcagggg ccaaaagaan ttcccaaaac cgaaaaatgat 780
 naaaaaaggg gactttgaa aaaaaaattt aaattttgc cagnaaagca tgggncccc 840
 cccttgaana aacccctgc atnaaaccaa ctttntggga ntttttngg tanggtttt 900
 ctggct 906

<210> 215

<211> 312

<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(312)

<223> n=A,T,C or G

<400> 215

ggcacgagga aaccaggtt gctgggtttt gggtgtaaac ttaaaaatga caatcagcat 60
 gagctggccg tggctgtgg gggtttagg ggcatttgg taagggaacc ctcgctcagt 120
 ccctctctgt tctgggggg aggacaagga gggcaatag gggccaatag ggaggctgct 180
 gctaggangg tttcctaaaa gaacaggtgt agggctaggg ctggttctt gttcaggttg 240
 ctctggcag tgatttatat ccacacaccc ttctgcaaag tgtcctaagg aganggcagg 300
 gataggatgt tc 312

<210> 216

<211> 341

<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(341)

<223> n=A,T,C or G

<400> 216

taaggctntc gaanataatg aatgagtcan ggagaggctn atgangaaat nccaaacacc 60
 tgactaatng gtgccacatg attncaatgg nctanacatg gtttagatct cntcngngga 120
 atgagcaata acacccntaa antcntcaat tgaccttagac acttcacact taaaanatca 180
 tcacttttaa ngaccacgaa tgatgcttaa gaatcacatt ttgtgnngaa ntggantctg 240
 gctacttaca cgaacagatt cttattcctg ttcatgagcc agtagacccg gaanaagact 300
 taagagcttc tgancttctt cttagctcca nngcttgaan g 341

<210> 217

<211> 273
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(273)
<223> n=A,T,C or G

<400> 217
nnccttcncc ccttnacnga catgaacaaa acagcngtct ngaaatttt aatacatnn 60
aagggttaacn ctcctnctt ntgtttccg nttaanncta nacctgcgcn gggcgccg 120
atncagccct atagttagaa gcctaattnc agcacactgg cggccgttac tanngnatcc 180
cgactcgta ncaantttg gngtaaagat ggacatanct ctatccnnga gnactcgta 240
nccnttcctt atnttacatg cnctaacgna gac 273

<210> 218
<211> 687
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(687)
<223> n=A,T,C or G

<400> 218
tttcagtg tttttgttc tcaatttga tgtcaaaatc tctgggtct tctaancng 60
ttatgttctt ccanaaaatc ttccagttt ttgttaattt tttctatatc agaagcgcct 120
ganccaaatg cccaaatnat acaccggct tctccggAAC gcttggcna aagggtntag 180
tcnatnngc tcctggaagc atctnaatg ctccaggtt ctcccangnc cctggannac 240
ttcanttgta tanacgaatc ctggtttgc agcggccctt gatatcgcaa gaaatacgg 300
taaaaattat ccaagctctc ttcccactna gganttcgga tctcatcagc cgggtaaagg 360
aaaactcctc angaagtttgc ggttccccct ccggcttacc ggctaattgtt aggaattact 420
tctggcttc ttccgataca tcctctctt aaagttaaga aggttaaaag aatnttaacn 480
tctcccagtgc gctaatggtc aaacaccatc ctcatnagtc agactgggtt ttcgaaagga 540
ggatataacc tccttgcna gttttaaa aaggattaa ccanatggac tanccctcnc 600
cccgaggattt nctcttcac aggagaagggt gtctcnccnc ttggctcatc cgaagcatag 660
gcaaaccnn gggaaattttc agaaacc 687

<210> 219
<211> 247
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(247)
<223> n=A,T,C or G

<400> 219
ggcccttcn cttttaatc gagagatcca aggttcaagg catgaaatac cagnctataa 60
aatgtctcaa gacntaaata atacggatng ngatagagag gttgaataat aaatgaanaa 120
anatgaaagn nattatgngg gaatacnaaa aaancngact aanggcggca ctgctggca 180
tggnnnaatc ggattaattc ctcataggac agccnaaccc cttaaatct canttccgt 240
nacccga 247

<210> 220
<211> 937
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(937)
<223> n=A,T,C or G

<400> 220
cgggctcgag tgcggccgca agctttttt actatagacc aatattaaag tcagtttaagt 60
tccaaataca ganttgaaa actaaagtaa aatattaat gggagaatat ctgcacatctga 120
atatgtcaac tgtttgctat tttagtgcata tttaatccctt ctacactgtat ctcagaaaca 180
aatttaaaaaa ttaatagatt tgacagcaaa atcattcagc accttactta ctccatcagc 240
aaggtaatata ttttagtcatt tccatccatg tggccaaact gaaaatccct aaccaccacc 300
aaccaaaaat aaataaaataa aaggagaggg ggtgggggga gagagagaga gaaagctcat 360
taaatagtaa aaaagtaaat aaaacaatga agttaaattc aggcctcagt aggcccagaa 420
actgtaaaca tttcacatgt aaatcatata caataaacac tgctaaaagt gtaaaattcta 480
ctggcttctg agatacaaatac acacgatgtt agggaaattct aagacatttc tacttggtt 540
atgcataattt aaaattcagg gaaatatcag ctattctacc tggaaatatgt ttaagaaaaa 600
ttccttatttt ctctaaaaaa aggaataatc agaagacgct acataactatg taagaaaaact 660
atacaatgac ccatcattag aagattcaga atagggaaaga aataataatt cactaataaa 720
atataatttat attgactgtc tttttttatg atagcaacaa tgattcagca taaagtaaaa 780
atataatgtat ttccgatgcc attttttattt cagttattct tttagtttc tgtagataata 840
attatctgcc tatctctgac ttctgancag tcatttatgt ccaattataa gtacatgtgc 900
atattttattt accttaaacaacg cctctcaat cctttca 937

<210> 221
<211> 353
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(353)
<223> n=A,T,C or G

<400> 221
ggctatnnna tnnttnaan atcntgnncnn ctttgacgct gttantaaan aaaaacaaac 60
gaatatcctt tttttgctcc cccctgtnc aataacttc tcacactaat acttacagta 120
taactnttcc tttcaactac caatattaag ttccaagcca cctgggctta agtatccaa 180
caacttaggt aatttggc taaccaccaat actatatgct aattataaca ctctaagccc 240
caaggaattt ttgttcagat ttcttatant ttccacttat aatatnatt ccncctctat 300
gggtattnn nnccctctagn cccatatnnc ccacngggat ttgttgaggg ggc 353

<210> 222
<211> 813
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(813)

<223> n=A,T,C or G

<400> 222

ggcacgaggc tttactaagg ccagactcac tatccccgt tctgttctgt ggtacactgt 60
tcactccca gtccatccca acctgacttc ctggccactg cagctttcc gataagggtc 120
agcagtggct tagttattgc taaataataa gcgcacatgc actcccttcc tcctgaaaca 180
ttgtccctcc ttggtttctg ttccctccca ggtctccat cactcccttct tagtcttctg 240
tgccggacttc tgttccttctt gcccattaaa agttggatt ttccaggatt ctgtccctagg 300
cccacttact tctcattctg cacgttcttgc ttggatgatt ctatcacatc cctaacttct 360
gctgccagt atgcacttaa aattcccaa tctgtatatac tgatctggc ctgtgtctct 420
agcctagaag tgtctttat cccagaagca cctcaaacac tgcaactttgg aaattaagct 480
tactgagtct cgagtctcaa gtcccaaact gacttcttt tctctatttt ggtagtgac 540
aacactattt attcagtcattt gcaaaccaga gcccgtgagaa ccatcttaca ttctcttct 600
cccttactc agttcttgc tctgttctt ctccctcncc tctcctgcct gtgggcctag 660
nggnccattaa ctgggttggca ctgcattact ttcnattttt ttggctganc taaccnnaag 720
ancctnttgt aggggcctt ctntcaggcn tnacttctnn caagancccc cgaaaccaga 780
tccnggggan tgctatggnn tggaaatattt ttg 813

<210> 223

<211> 882

<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(882)

<223> n=A,T,C or G

<400> 223

tcacactact gagaaggcagg gaaacccact gaaaggcac gtttcttaac ctcagaatgg 60
ggctactagc ctctaaagca ggaattgegt tttgttttagt atttccatgg tctgctgcaa 120
ggcgtggcct ttacccaatg gataaatgcg tacaaggctc ttgtgagcag tcaagttct 180
cgaggtttac agttgaaggg aagtgggatt gtttccctgc gcatttaat gaaggttaggt 240
gggtgatcac ctttccttaa atgtgtgaag ggatgagata aagagatagg catcttaatt 300
gccactgtatg gccttcaggt gaggacaggc atgagccaa tgaagctttg acaattgtgc 360
tgaacccaaa acttcaaaaaa caagaaaaaa catagactgg ctgaaatgat ctaagtcaac 420
agagcatggc cagcgcttca tacaaggcag gaccacaggc gaacactgac agcccaggag 480
gcactgagac agaggcagtg ggaagaagtg acagacccca gggactcccc accaacagca 540
gctgctgttgc attaggaacc cccagtagac tgtcaggcac ctggtagtgg agaggctacc 600
aaggcccgga ctggagagga gccaaaggaa gaaacagtgc agtgcttaga cccctctggg 660
tctgcccgtg tccataccccc tagggagatt ccattccaga agtggacata ttcccacaga 720
gtgcctgggg ctcaactcatc acagctgccc ctncatgaag gcattctcac tgcagccctta 780
ncagggaaca gggtcatttg cattaggcan cttgctgtcc tagaaggcn tggngntccc 840
tacactgccc atgttcccaa ngngttcaa nctcnaaaan tn 882

<210> 224

<211> 660

<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(660)

<223> n=A,T,C or G

<400> 224

gattaaactc aatcattcac ccgggctcga gtgcggccgc aagctttttt tttttttttt 60
ttttttttt ttttggncct ctgggcttgt gcccggaaagg ggantgctgg gccacntggg 120
tgtccgtgtt tgatttctg ggacctgccc ccccgtncc cgccccggnt gccgcgtctc 180
actccccgcc gcggtgenag gggccccgtg tgccgcac ccttccaccc gtgtttgt 240
gttttttga ctntggcgt cccaggggtg cancggccgt gggggccctgg tttgctttca 300
cctcttcata tgctcaactgg ccgnantgn gtcttnnca aacaaacgtn tgaaggncaa 360
nccctggct cctgtgaacc cggccgtt tgcggcaan tctgaggctc cttcgattt 420
ctggatccgg cctntggctg gangcgtgt ctgcaggcac tgctcccatt gctggcancc 480
ttttctcccc gtggccgccc ggcgcggcat naaggcggt gcaaacgccc gccctgcca 540
gcgcaaagtc aaacncgggt gcggcggga ccccccggcg gnccggaaaca cccancagg 600
cgggcaccac aanaagcgcg gnccctccggc gtctaaaact nccatgtggc nccccccgn 660

<210> 225
<211> 438
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(438)
<223> n=A,T,C or G

<400> 225
aaaaaaaaaaag gaaaagtacc cagtgtctc agcttcttag cctcctctac agccctgttg 60
gnntttaaac ctgtgcctg tgtctgtgtc cccacttaat atatatagta cacagctgga 120
gagatggctc agccaggaga gggaccata ggtctgtgaa ttccagagga naggcaggna 180
tttataggtg gntctgtcag gtgaaatcng aggagccaa gctattgtat gtcatatgt 240
cagccggct ctgtgggagg tggtgtaaga cctatgnat gggacangtg tncacgctgg 300
gatctctggc cggttccgaa aagtgggat caggtagtgg gtggctgatt gcacaagttt 360
anaaccagg attaggacac cacaggtcag cacctgctc tcagcatcct gactgggtgt 420
gatggcata ctcaaggc 438

<210> 226
<211> 480
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(480)
<223> n=A,T,C or G

<400> 226
aaaataaaaa caaaaaggat ctttagaggc ctttacttca gtggttctca atgtcagagg 60
atgttatgt acctaataa aatctccagg ggaactgttt tgaactcaac agactctctc 120
ctgttctgag agactctgac aaagttggga gagctgcccag gtactgtcca catgaccctg 180
actgccatg attcaattac cttgaatggc ttatccagtc caataccctc atttcttaca 240
tgagggaaact gaagcacgta tcacatagt atacaatgaa aacttggct taatcgattt 300
tcagtgctgc cagtacaatg tcttgagcat atcaatttct tccaaaccctt gacaacataa 360
ggtagcaccat ctaaaattttt tatttctgct aatttattag accaaaaaaaaa aagggnatct 420
cncccatgtt ttacaggga tgattttatt ncagaggatt tcacntggn gctgattcnt 480

<210> 227
<211> 423
<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1) ... (423)

<223> n=A,T,C or G

<400> 227

cattgttgtg ggctctgctt agcacatcac atcggagcac agaggtgacc tttctgcc 60
cagggatgtt caccttagtc acctgattga ttcccttca cttggcac gtgattcctc 120
caggaggatg ttcaccttg tcgcctgatt cctccaggag gatgttcacc ttggtcgcct 180
gaccacacag gcatctatca ggcttctca ctgcagccac tatgtccccaa taatggatga 240
gtgtcttgtg gagagatagt ccaaatacaca ctgataacctt ttgcctcata cggcctcacc 300
ccccaaacaat cnaccactaa tgactgcctc atagcagttt ttccatttcc acagttcctt 360
ctatatgtat taattgtcat tctactataa agaanactt ttctttaaa aaaaaaaaaa 420
aag 423

<210> 228

<211> 249

<212> DNA

<213> Homo sapiens

<400> 228

cattgttgtg ggctgttagta aaatatgtgt ctggtaagat atgtgaagaa ataaaataag 60
atcaattaaa tctggcccat tgaatgacac attaattgtt tattaatatg taatgttaaa 120
gatatttagga gatggtgaaa cattatggca aactaaattt gggaggaggt tgaattgtat 180
aatttatgaa atccctaaagt ctgtacatt aacactctt actgtcaact tttcaaagca 240
gtgagaaac 249

<210> 229

<211> 436

<212> DNA

<213> Homo sapiens

<400> 229

cattgttgtg ggatgttatac tgaccatcac aatatgattt ataatatggaa ggcataagt 60
catttctcat tggggcagga gtgtggcaag gggaaagaag agctttacca attaactcaa 120
gattatttgg tgacatttctt cttacctttt aggtgaggag aaagagacag aggatggaga 180
attggtgtctt ttagttagct gatacattaa gctgcctgga agcagatgtt aaatcctatt 240
gaaaataatt ttatgtcgat tttgttttgcgt gcattgttta gcaaaataact acacaaaaag 300
tcttgacctg tgggtttgaa atggcagatg ttcacagtg ggactgagcc ttggggcaac 360
atcaatcttc acaattctgc acctatttgc tcaataactg gcttgggtgg aaaaaaagg 420
aaaaaaaaaa aaaaag 436

<210> 230

<211> 760

<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1) ... (760)

<223> n=A,T,C or G

<400> 230
cattgtgtt ggnngtgaa ggaaaanttt gaggcaatga agctaaacat aaaagaggaa 60
aagcanatgt taccta atg accacaatct acaaagtcca aatanaaaac ctgggagtt 120
gataggatga aactataacc tccagcaaag agcttaacag caattaaaat aaagacaaat 180
ttctggatg gatnagacaa agtagcatat attacaaagg aaaatanaact agtatcatnt 240
acgtttgatt aagtaactgc tttcaaataa ttgaatcata aacaatgatt tctgcggtt 300
taagctcatt atttggttc cctggttct cctaggatgc agtataagaat ctccatgcct 360
gatgtttatg taccacacaga agctgctgct tcttttttc attatttcct ttttaagtga 420
aagtttaatac cttttatatg ttacagagaa gaggcagaaa aagccacact cccactatgc 480
tattaaatgc cctgaggatc aactgaggaa tgattatacn catggctgaa tacagtntat 540
tcatttgttt ctttggatgt tanataacaa aaggtggat tctgtaacat cttgtgncaa 600
ttanccaaat gttaaggcga aaatggaatc tttcaaacaa gtgtntaaa caggtttga 660
tttccaaaaa ttantatta gaaccntttc aattctggaa gttnccaa at ttccangttg 720
tgtttctct tccaatttctt ctccctttg naaattcccc 760

<210> 231
<211> 692
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(692)
<223> n=A,T,C or G

<400> 231
cattgtgtt ggggggtgctn tggggagaac acgcttatgt tganatnggg ctccccgaga 60
aaggcatttgc gacacnttcg aataaggacc cntngggaaa ttcangttag ttgtggacat 120
ncntagataa natcaaaggc cttgangaaag tccgcctggc accttccngt ctgcgaggag 180
gttgatcca aatgctaagg ggtccagntg cantgtanta tcgtgagatc agagtgtatgg 240
gcaggtgtgg gcatgcgggc cctcaanang aagtgcggcag gatgactcag acttatgcct 300
atatccattc antcctgttc attatttttta ncntccctc naaggacccc caattnaac 360
catttttat tcanggctat acttataaaa gtcattgtt ttnagtcgg gtgatattaa 420
aaccatttgg acgccangca tggggctcn nggcctataa tccntccac cttggggaaag 480
ccgaagctgg tnnaatccct naaggtcnngg aatttggaaaa ccattcctggg ncaacattgg 540
gngaaaccct gtctctactn caaaaaacan aaaattttct ggggcctngg ttnaggtgn 600
gcctgaaaat ttccancnt tactccggaa aggccgaatg ccntaaaaaa nnacctta 660
accccccga anggcggaa agtttccatt tn 692

<210> 232
<211> 518
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(518)
<223> n=A,T,C or G

<400> 232
actcaaatgn ccncttgaag gtcacccaga ctcanaaagt gtcaagctt ggggtgggt 60
gtaatnaata nctcggnctc ctgattagtn ctcctagctc gatenctggc tgagatnngt 120
tcgagcaccc ttcctttgtat cccgtcaaac nccnggnaaa agcngcctgc gtagtcncct 180
nagccgaatc tgnntttcccg acaccctccg ctgcgtcggc tgccctggtn aagcngcnc 240
ctnaaanaan aaagngaagt ctccccngtc tcncccnant cctngggaaa acngcctgaa 300
ccaatatgnt ccccccaaggn cnccccaggg cacntaaccc gtttaggaggg ccccccnc 360

gcgtttggc cnnaagccn gccccngnaa taacccnct anaaccacgn aaaaatgcaa 420
agtccaaag gtaaagaat ctcccnaccc cccggtccc tcgcaanctt cccctnnngna 480
cttgttcc gggaaaaccc ttanccgan ccttcga 518

<210> 233
<211> 698
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(698)
<223> n=A,T,C or G

<400> 233
gcacgagttt ctgtctgtct gtctctctct ctctctctgt ctctctctca 60
cagttagaat ttggctgtt tctttattca atacccaaat atatgttcat tagggtata 120
ctgtatacac tacacataac agttttgttt tttgtttgg atattatttg ataataagaa 180
ttttaccaca tcattaaaaa aagtttcccc aagctataat tttgataat tgcaactttc 240
cactattcaa atgtttatTT aactcttct ctcctggagt aggtttacat tccatTTtag 300
ctatgatact gcttaagag aaattgtttt aagataaatt tccatagaca ggtcaaagga 360
ggtaaatata tgtaagctt tcgatgcctg ttactgaatc tcattctgga aaacataact 420
gtcaatgccc tcttttctc atggtaaaaaa aatacataac aaaatttacc atcttaatcg 480
tttttaatg ttacagtacg atagtgttna ctgtatgtac cttgtgcaac agattcttg 540
aaaactttt cattttcaa aatgaaaact ctgtactcat tgaacaggca gcttcccaac 600
ttccccattc ctcccannc ctacccctgg ttaanagtct nacaaaaccc gggaaTTta 660
tgaaatttga aacacttttta naataccnnc tattaggg 698

<210> 234
<211> 773
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(773)
<223> n=A,T,C or G

<400> 234
ggcacgagcg cagttttcg aaagctgtaa tttgtttgt atcaaaagtc ctgcagtata 60
ttagtcctat tgcattttaa agagttcca agtgcattgt gatgggtgtc tgTTTTtag 120
tattacggtc ttatgtatg ttgcAAAact agtcatttg gtgtgtcgt acggggcgga 180
aagatcaggc caggcaaaat actctggccg ccaaagtaaa tgcttaaggc cgccaaacgga 240
ttatgtcctg gggttcgatg agggccgtaa ttaggttgag ctgggtgtang ctaacctcgc 300
agccatgtcg gagagagatg agagacataa natttaaag taggggcgtt ttttacgaag 360
ttctgancca tttcTTTGT tatcggtccc ggcaaaagca actgagataa atgtttaaa 420
agactcgatg attttttcga cttcagcaac gtactcagcc ttgggttctc gtatTTTC 480
aaaggcagct atttgctgag attcatgaaa agtttgactt ganctgctt tcaatttctg 540
cagcncgggc ttcaactgtt attgaatttgc ttgtattaag cncaatacgt tgcnngtcac 600
caaggTTTC catgtttga ctncacctgg tcgaaccaat ttgaattatg tnttttgcc 660
tgnccgttc cccncctt aaatccatct ctttttnga aacctttngn nggttgaatt 720
cngccgcccc gttcccaacn tttggttcna ctttggaaaaaa aaanatgggt agt 773

<210> 235
<211> 849

<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(849)
<223> n=A,T,C or G

<400> 235

attgggtacg ggcccccctc gagcagcctc cactgcaatg ccgctgaatc aagagacttt 60
tcaatacgct ttatcagtga aaatgatgtg atctgaagag tcctatcttgc agcactttgc 120
atgacatcca acgttaatgt ccacaacgtt ctttagctgcc caacccttt atcggcaagc 180
tccaaagggtg ttttgcaaacg ttctacggcg tcatgaaaag ctgaaaaatg ctgtgtcaac 240
actgcaccgc tgccatctt caaaagcagc gcccttatag tctccgcatt cgaagacgt 300
aaccgcgtt gaatagcctc ataatcactt ttgttagaaat caatcagagc tgtgcttagga 360
acctttccat cccaaacata cgactgtgc accacgtctg caaaaggcaga cgtcacatta 420
tgcataatgcc ctcttaccgt cagccgatca tcctcactca tagcgcacgcg agaaagctct 480
tgttccagct cgtgcacggt atccaaattca gtaatectac gcaacgcgt ctgaatcg 540
ttcataagtt cagtttaaa gctcaaaact tcgtcttta nttaaaaaatc tgtgactttc 600
aaactggcg antcttcacc attttattaa tcgtctttt gangganggc ccagcgtag 660
atctgcattcg ccagcgaaat cgttactccc tcccattctt cctccggta acgcanntag 720
tttctccgaa gccttaaaat tagccgggaa aagggaantt atttgccttca acaangnat 780
cgccgnctg gtggtaaaaa ggaactgaaa taaaattaaa nccncttgg gggaaangcc 840
cgccatactg 849

<210> 236
<211> 310
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(310)
<223> n=A,T,C or G

<400> 236

gggggtgggtt gcttccgaaa nccggggccc ggccaacttg ttggcttggg aatattctgg 60
caagaaaaatt tccagggcg cgccaaattttn atcaagcccc ggccgcctta aaccgaaaaac 120
tctggcaggg tcaaccctt tcattggcgn ttgaaagctt gaagcgcccc aagttactcc 180
caagcttggcgtt gcgnttgcgg ttggggcg ggaaaaagtt gaaaacacgg gcgnttggc 240
gcccgcggcccg cgggcggttt nttacgcccatt cctggaaaaa ctttcagggt tggctgctt 300
cnaaaaacggg 310

<210> 237
<211> 315
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(315)
<223> n=A,T,C or G

<400> 237

gcacgagtnt ttgttattta natnttgctt tggttaangg aagaacacaa naatgcctg 60
ctaaaggat tctgtttqqt tgcanqctgc naqcqqqqa aaaatcnaan tqtatnttqc 120

acaacangat ttttagaan tcagaactat gacatgaagt canncagggc actctacgac 180
tgaatttgcn gtgctgcctt cacangctcc ttntcgctc tntnctggca ncngtgactc 240
ntacacgtcc tgganantan cctccctana aggaacgact ccgacacccc cccnntaccc 300
ctnaangttc atcng 315

<210> 238
<211> 510
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(510)
<223> n=A,T,C or G

<400> 238
ngcacagtn ttttttattt atatattgct ttgtttaaag gaagaacaca aaaatgccct 60
gctaaaggga ttctgtttgg ttgcaggctg cnngcgggga aaaaatcaaa gtgtatTTT 120
cagaaaaatga ttttttanaa gtcagaacta tgacatgaag tcaagcaggg cactctagga 180
ctgaatttgc tgtgctgcct tcatatgctc ctgcgtcgct ctttctggc agctgtgact 240
cncacaggtc atggaganta tcattcccta aaaggaacaa cnccgatatt catcttatac 300
cattaagtnc atctgtccca ttctatgtng tggatgctaa ctttgatca ttgatngtga 360
tnccatggac atntancatc anctttcana ncctnggatc tttgacnagt cttattantn 420
agantccaac tantacgtg ccganttana aatgctggnt ntccaaattcc tactcaaata 480
nccnacatga acttccantc ccctgcnnna 510

<210> 239
<211> 209
<212> DNA
<213> Homo sapiens

<400> 239
ggtgcttttc ctttctactc gtcttcctgc ctggcaggag aagctccgc tactggttgc 60
ccttctacca ctgtcgacac caccactgc agtgagccag tgtccgaggc tccagccaga 120
aacaggtagc agccatgcgc gataccaaac gcccacactt aagagcctga aatgacctga 180
cgccacctcc gcatgcttta cctactgag 209

<210> 240
<211> 610
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(610)
<223> n=A,T,C or G

<400> 240
ggcacgaggt ttctggctgg agcctcgac actggctcac tgcagttgggt ggtgtcgaca 60
gtggtagag ggcaaccagt aacgggagct tctcctgcca ggcaggaaga cgagtagaaag 120
ggagcggcat gctggaggt ggagcctgag cccctggggc tcgccttgcgt gtgtttgggt 180
gtgacgtggg acactgcagc tcggccagag tggtaaaaaa tgcctgggt tacgctttc 240
tggcttgcg cgtctatctg ctccaaagcca ggctgganga ngagganaag gaatcacctg 300
tggtacgctg gagcctgcat gtggcgtgac tctgcaactc gcctcgatgtg actgatggca 360
gccacggaga ctgcagctcg acagggagtg aggcttctca ntggcttgaag 420

actccccacga aatttgcggg aaactcaagg ctgtcagtga ctttcgtggc gccaagactt 480
aancangcgc gttgcattgca tccggccagt gtctgtgcca cgtgccctga cnccaccttg 540
anataancac ccggAACGCG cnncgcgcag gccgcgcgca cacgnccggg cancaacttg 600
gctggcttcc 610

<210> 241
<211> 474
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(474)
<223> n=A,T,C or G

<400> 241
ggcacgaggt ttctggctgg agcctcggac actggctcac tgcagttggg ggtgtcgaca 60
gtggtagag ggcaaccaa aacgggagct ttcctgcca ggcaggaaga cgantagaan 120
ggancggcat gctgganact ggancctgan cccctggggc tcccttgcgtg tgtttgggtgg 180
tgacgtggga cactgcagct cggccagant ggtaaaaatg tcctgggtgtc cgctttctg 240
gctttggcccg tctatctgtc ccaagccacg ctggaaagang agganaagga ntacacctgtg 300
gtacgcccga gcctgcatgt gggngtgaact ctgcaactcg cctcgtgtga ctgatggcac 360
ccacggacac tgccactcta cagnaatga ggcttctccn tggactngaa agctcanctt 420
nactccncc aagtttgncc gaactcaagg ctntcaactna acttcgtggc gcca 474

<210> 242
<211> 415
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(415)
<223> n=A,T,C or G

<400> 242
ngcgggnnt tccaccagct cgtgtgcaca agtngcgcca cacaacatg cgcaggcact 60
gcatgtcata natgtgcttc gccgtggttc tggaacagcg agtagaaatg ggcgttcggg 120
tcgcgaccaa attcgacgtc ntggatgctc ttgcgcaga angtcacgtc cgggatcggc 180
ccgatggatc cgctnaagcg ccgaaaggcc ctgacttgca aaccgcggct cacagaaccg 240
gcaccaccgg cgccctccgc cnacaaaatg cgagcggctt ccgacacacaca ctccctcaca 300
tccccgtcnc gcacttcggc ngtttctagc tccgcccacgg ttgtcagcgg caccgcggc 360
gccnagctgc cggcggcata cgttgcacac agcacacacg gatccgtct cgtgc 415

<210> 243
<211> 841
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(841)
<223> n=A,T,C or G

<400> 243

aacgagggtgt cgatgagcgc gaacaatcgc cctccttcat ctctacctga tggtaactt 60
cgctcctaca gcccggccaa tgaagacgaa tggctgctgc cgaggatgg agtctca 120
gagcacgcgg cgctggacaa ctcatcgact ttgtacgcttc cggttagctt gcccattcag 180
ctccactgac gacagagacg gagctggcca ctgccatctc gacgcagcgg gacaaggagc 240
agcttcgggc gccgtatgca tcactcgaag agaaccagga gcagccggaa gcaggangcg 300
ctgcacggta caggcactt cggcgttca gcgatccat cggccgcattt ccgtacgtca 360
ccttcttgcg caagaacatc caggacgtcg aattcggtgc cgaaccgaat gccatcttct 420
actcgcttcc caggaccgc gcaaggcaca ttgtatgacat gcagtgcctt ggcgtatgttt 480
gtgcggcgct accttggtgc acacgaacga nggcaaccaa cccgcggcc 540
atgcatttctt gttctgttcc ggtgtgcattt gccggatgtt gaccgtganc ttggtaatc 600
ggctggtgca tgaagactta ccgcctctntt caagggcgaa cgcncctcan ttgganaag 660
gaacaaaacc cccccnnaag aacggcantt gcancntttt ccccgctgc cggctttctt 720
ccattcgggn attctctntc tccnaaaantt ccgcnaaatc ttcttcggg ttctccctg 780
tttttatttg cccttccgc cacttgggtt gtttacatc ctacaancct ttttttctc 840
c 841

<210> 244
<211> 761
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(761)
<223> n=A,T,C or G

<400> 244
aacgagggtgt cgatgagcgc gaacaatcgc cctccttcat ctctacctga tggtaactt 60
cgctcctaca gcccggccaa tgaagacgaa tggctgctgc cgaggatgg agtctca 120
agagcacgcgcg cgctggacaa actcatcgac ttgtacgctt ccgttagctt agcccattca 180
gctccactga cgacagagac ggagctggcc actgccatctc gacgcagcgg ggacaaggag 240
cancttcggg cgccgtatgca atcactcga gagaaccagg agcagccggaa agcaggaggc 300
gctgcacggc acaggcactt tcggcgttcc agcggatcca tcggccgat cccgtacgtc 360
accttcttgc gcaagaaaaca tccaggacgt cgaattcggt cgcgaccgcg atgcctatctt 420
ctactcgctc ttccaggacc cggcgaagca catttgcatttactgcgtgc ctgcgtatgt 480
ttgttgcggc gcatacctggt tgcaacncgan cganggaac aaccgcgcg angttgcgc 540
tctatgcatt ccctgtctgt ccgggtttgc atggccggat gtggancgtg ancttgtaa 600
tccgctgggt gcatgaagga cttaccgctc tcgtcaaggg cgaacgcgcg atcaattccg 660
aaaaaggaaac naaaaccccc ccccaangac ggnatttgc anctttccc ncncctgcg 720
gctcttctcc antncgggct tctcttctc anaaaaattcc c 761

<210> 245
<211> 710
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(710)
<223> n=A,T,C or G

<400> 245
aacgagggtgt cgatgagcgc gaacaatcgc cctccttcat ctctacctga tggtaactt 60
cgctcctaca gcccggccaa tgaagacgaa tggctgctgc cgaggatgg agtctca 120
agagcacgcgcg cgctggacaa actcatcgac ttgtacgctt ccgttagctt agcccattca 180
gctccactga cgacagagac ggagctggcc actgccatctc gacgcagcgg ggacaaggag 240

cagttcggg cgccgtatgc atcaactcgaa gagaaccagg agcagccgga agcaggaggc 300
 gctgcacggt acaggcactt tcggcgcttc agcggatcca tcgggcccgt cccgtacgtc 360
 accttcttgc gcaagaacat ccaggacgtc aaattcggtc gcgaccgaat gccatcttct 420
 actcgctctt ccaggaaccg gcgaagcaca ttgataacat catgcctgcc catgtttgtt 480
 gcggccctcc tggttgcnca cgaancgaag ggcaacaaac cccgcggcagg tngccgctct 540
 tatgcattcc ttgtctgttc cggtnntgca tggcccggn nttggaaccc tnancgtt 600
 nnaatcggtt ggtgcattga aggaacttac cgctctcgta aaggccgaa cgcnccttc 660
 agttcggana aaggancgaa aaccccccna aaggaacgg ccntgcnng 710

<210> 246
 <211> 704
 <212> DNA
 <213> Homo sapiens

<220>
 <221> misc_feature
 <222> (1)...(704)
 <223> n=A,T,C or G

<400> 246
 aacgaggtgt cgatgagcgc gaacaatcgc ctccttcata ctctacactga tggtaactt 60
 cgctcctaca ggcgagccaa tgaanacgaa ntggctgctg ccgaggatgg gagtctact 120
 aaagcacgcg ggcgtggaca actcatcgac ttgtacgtt ccggtagctt agccattca 180
 gctccactga cgacaganac ggagctggcc actgcatct cgacgcagcg ggacaaggga 240
 gcagcttcgg ggcgttatg catactcgta agagaacagg agcagccgga agcaggaggc 300
 gctgcccgtt acaggcactt tcggcgcttc ancggatcca tcgggcccgt cccgtacgtc 360
 accttcttgc gcaanaacat ccaggacgtc gaattcggtc ggcacccgaa ttgcatttt 420
 ctactcgctc ttccaggac cggcgaagca cattgatnaa attgcattgc ctgcgcattt 480
 ttgtgcgggg cttcttggtt ccccgancga agggcnacaa ccccgccca ggggccnct 540
 ctatgcattc ctntctgttc cggtgttgcn tgggcggat ttgaaccgtt aancttgggt 600
 aatccgnttg gtgcattaag aacntaaccg ttcnctgtca ggggcnnacc ggncccttnc 660
 aatttcggaa aaangaacca aaancccccna cnccaagga aacn 704

<210> 247
 <211> 618
 <212> DNA
 <213> Homo sapiens

<220>
 <221> misc_feature
 <222> (1)...(618)
 <223> n=A,T,C or G

<400> 247
 ggccgcccagt gtgatggata tcgaattcaa cgagggtgtcg atgagcgcga acaatcgccc 60
 tccttcatact ctacctgtat gtgaacttcg ctctacagc cgagccaatg aagacgaagt 120
 ggctgctgcc gaggatggga gtctactag agcacgcggc gctggacaac tcacgtactt 180
 gtacgttcc ggtagcttag cccattcagc tccactgacg acagagacgg agctggccac 240
 tgccatctcg acgcagcggg acaaggagca gcttcggcg ccgtatgcat cactcgaaga 300
 gaaccaggaa gcagccggaa gcaggaggcg ctgcacggta caggcacttt cggcgcttca 360
 gcggatccat cggccgatc ccgtacgtca cttcttgctc caagaacatc caggacgtcg 420
 aattcggtcg cgacccgaat gccatcttct actcgcttcc ccaaggacccg gcaaaagcac 480
 attgatgaca tgcagtgcct gcgcatttt gtngcggcgc tacctgggtgc acacgagcga 540
 nggcaacaaa cccgcggccca ggtgcggctc tatgcattcc tggctgtcc ggggtgtcat 600
 ggcggatg tggaaaccc 618

<210> 248
<211> 622
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(622)
<223> n=A,T,C or G

<400> 248
gcacgagagc ggatccgtgt gtgctgtgt caacggatgc cgccggcagc ttggcgccc 60
cggtggcgt gacaaccgtg gcggagctag aaactgccga agtgcgcgac gggatgtga 120
gggagtgtgt gtcggaggcc gctcgacttt tggcgccga gggcgccgtt ggtggcggtt 180
ctgtgagccg cggtttcaa gtcaggccct ttcggcgctt cagcggatcc atcggccga 240
tcccgtacgt gacccctctt cgcggagca tccacnacgt cgaatttggt cgccaaaccga 300
acgcacatctt ctactcgctc ttccagaacc cggcgaagca cattgacaac atgcnnntgcc 360
tgcgcattttt tggcgccgc tncctgntgc acacgaccga gggatccaac ccgcgccagg 420
ntgcnnctctt acgcattctt gtcggccgg tggcgctggc cnggatgtgg accntgagcn 480
ggngantccg ctggtgcnntg aagacnttgc cgctctcgctc aaggccnacc gcccnnctcg 540
gcggaaaaaag gancaaaaanc ccccccgc当地 gaaccggcnc tgcaccgttn tcgcgcccc 600
gctgggctct ttcnnnttac gg 622

<210> 249
<211> 517
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(517)
<223> n=A,T,C or G

<400> 249
cattcgagct cggtagccgg gatccgattt gtaaaggggg tgcggaaacag ccagctgg 60
tttcgggtgc ggccggggca gcccacatcg ctgtggcgt tgccgtactg gatgcgtgt 120
gccgggacaa acgcgttttc caccacgatg tcatgactgc ctgtgccgcg caggcccagc 180
acatcccaact tggctcaat gcggttagtcc gccttggca ccagaaaagt cacatgtcc 240
aggccaggcg tgccatcacg cttgggcagc agaccgccta gaaacagccca gtcgcaatgc 300
ttggagccgg tggaaaagct ccagcgaccg ttgaacctga atccgccttc cacgggctcg 360
gccttgccag taggcatata ggtcgaggcg atgcgcacgc cgttatcctt gccccacaca 420
tcctgctggg cctggtcggg gaaaaancgc cagctgc当地 ggggtgaacg ccgaccaccc 480
cgtaaatcca ggccgtggac atgcagccct ttaccaa 517

<210> 250
<211> 215
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(215)
<223> n=A,T,C or G

<400> 250
nntncattgg gccgacgtcg catgctcccg gcccgcattgg ccgcgggatt accgcttgtg 60
accgcttgtg accgcttgtg accgcttgtg accgcttgtg accgcttgtg accgcttgtg 120
accgcttgtg accgcttgtg accgcttgtg accgcttgtg accgcttgtg accgcttgtg 180
accgcttgtg acngggggtg tctggggac tatga 215

<210> 251
<211> 231
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(231)
<223> n=A,T,C or G

<400> 251
ngcgcccacc tngtattttt tggtcgatca ctatcaagta tgtacatctt gctctagaca 60
actccnattc agtggaaagaa attggaaag tatccggat aagtaatagg nattaggct 120
nccttantgc ttggtggat attccncaac tgntccngat cgatcagnnc tcgtgtcn 180
aatgtgctc gatcgtnatt ctactnctga gcttctatcc nnacgtggcc t 231

<210> 252
<211> 389
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(389)
<223> n=A,T,C or G

<400> 252
atgtatcanc nctgttggtg ttncatcttt tgcatcgatcngt tctaaggcn gataantatc 60
agagatgcta atgcatnttc tgccaggcca ncattgggtgg cctatgcgtt ctcttcttat 120
cttcctgaag agtcatctt ggnggatgtg ttccccccctc tccacagtgt ttgcaagcgt 180
tacccacgcn tgtcggngcc gggaaaggtn ncacatccgg gnagacttcc ccncgtntga 240
atcgtnctn gaatctccgg cgtcntccct naacctctt actnggacaa ngnccgtnt 300
tccccctntgt gaactngtan cccgccttccct ttccccccctc agcctaanc ggaangaaga 360
cngggtcnat ctngggcncc acaagaant 389

<210> 253
<211> 289
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(289)
<223> n=A,T,C or G

<400> 253
ngggccnna tgagcgcgtaatacncatc actatngggc gaattggta cggggccccc 60
tcnagcggcc gccttttntt nttttttntt nttttttntt caaaacaccc tccnccntgg 120
atgganacgt nacccttctc taaccanatc ttcacaatnc nantctcagg cagccgcctc 180

aaanccgatg tcangttggn atntcaantn caatcttatt ttgngaatta anctganatt 240
gtggatggtn naccaatcan atacttggna tccgttgaac ccctgtgga 289

<210> 254
<211> 410
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(410)
<223> n=A,T,C or G

<400> 254
attgtgtgg gaactttag acagctatac caattgcagt gctatttctc tgaggtattg 60
aatctcantt attataattt tgaaatccaa ttggcttggaa ctttcatttattt ttccaactaa 120
aaagatgattt gaaggatttta tttgaaatgt gtaaagagta atatagattt tatgctttag 180
tttccttgaa aaaagtaggtt aaaattcttc tggaagtgtt actcctaaaaa tacaaatgaa 240
catgtcaaga attacataaa ttctttaaac tateccttaan aannaatggc tctatgtann 300
gagngaccct tacagactat taagaattaa cttgcatttgc anagactcat ttanattcat 360
gaaatgntc tcactttctt ggtaagatct ggcttggacg tttttggtaa 410

<210> 255
<211> 668
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(668)
<223> n=A,T,C or G

<400> 255
ttttttttt ttttcctgtg ccaggcacta taccactgtg ctaggtgcct tctttgcatt 60
acttcatttc ctcataagct ttctgaggan acagaaaagct tgaggttcac gtagctagca 120
tctacataaa tttagttgcta aaaacataca atacgttttc cggcaggctg tcatttagtaa 180
ctgatactac tagttgataa tctcataaac ctgcanaan ctaccattta agctgaaaca 240
actgtcaata tcaactaanta aaacttaaat ccataaatca actatattct aaaatctgac 300
ttcagttcaa ttaaaaaatc actagttgtt acctacctcc ttctgaaagc cagtacaagt 360
taaatgaaca actcccgagt ttaacaaaca agtggcatct aaaaaaaaaa tttaaaaaat 420
aatccactta catatattta aaatggcatt aataaaaacaa aatttatcca ataacnaant 480
ggcaaaggaa ggtgtccaat tattacatgt tataaatctt taaattaaac ttttcttngg 540
tttttcttcc ctanaataaa tacaancctt tccccgcna accagaaaaa agcaaaaaac 600
aaaacccaaa aactcccagc ncngcttaaa aaacncaaaa aaaataaaaan ctcttattaa 660
tgcccnaa 668

<210> 256
<211> 487
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(487)
<223> n=A,T,C or G

<400> 256

cgnaaccgtn cnttttnat gtgcgccgc cncagnacca gngccgctac aggcgaaggc 60
cggaagcacg ggagaggntt ngaaaaaaa agagtgccta caaagagcat attcgagag 120
ttggatgag tgaaggggac cagaaggngc acggtaggg acgcgtaaa ggangcneg 180
gagaatgac agcaagaagg gganaagcac acgaaaaggc agtatcctcc tcccccttt 240
tcgaggactg ccgcacatccc gtttctgcc cattccagtc accgaanaag atccaaana 300
aagaagaaaa gaancagagg tgcacttcgc ttcatatttc ntcgcttc tttctgnct 360
tcacnagttc tgcaggattt cccttgctt ctccgagca catctacgca cgnatgaggc 420
tcggcaggta aagccnacaa aacnctcgca ctcctttt tcttgcnnng tctgngtggt 480
anggnng 487

<210> 257

<211> 502

<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(502)

<223> n=A,T,C or G

<400> 257

ccttggaaag nccngctnaa ttcngnganc cccngatca gcaccaggga gctacaacna 60
aggccggaag caggggattt ngccggaaaa aaaagagtgc ttacaaagag nttatccnca 120
nagatggat gagtgaaggg gacgagaagg tgcagcgta gggacgcgtg aaaggaggca 180
gcggagaaat gacagcaaga aggggagaag cacacgaaaa ggcagtatcc tcctcccccc 240
ttttcgagga ctgccgcattc ttttttctt gcccattcca gtcaccgaaa aagatccaa 300
agaaaagaaga aaagaaacag aggtgcactt cgcttcatat ttcgctcgat ttctttctg 360
tcttcacaag tctgcaggat tgcccttgcc ctctccgag cacatctacg cacgtatgag 420
gctcgagggn caagccaaaa aaacgcttgc actcctttt ttcttgcgt gtctgtgtgt 480
atgtgaaattt ccgcggcncc gc 502

<210> 258

<211> 510

<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(510)

<223> n=A,T,C or G

<400> 258

actcgncact cgatncanta caagagnnta tgnattcgaa ngtccccccg catcagcacc 60
agggagctac aacgaaggcc ggaagcaggg gagagggccg gaaaaaaaaa agtgcattaca 120
aagagcatat ccgcagagtt gggatgagtg aaggggacga gaaggtgcag cgtagggac 180
gcgtgaaagg aggcagcgga gaaatgacag caagaagggg agaagcacac gaaaaggcag 240
tatcccttc ccccccttgc gaggactgccc gcatcttgc tttctgccc ttccagtcac 300
cgaaaaagat cccaaagaaaa gaanaaaaaga aacagaggta cacttcgctt catatttcgc 360
tcgcttctt ttctgtcttc caagtctgca ggattgcctt tgtctcttc cgagcacatc 420
tacgcacgta tgaagctcg aggtcnngnc aaaaaaacgc ttgcactcctt ctttttctt 480
gcnagtctgt gtgcattggg gaaatnctna 510

<210> 259

<211> 292
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(292)
<223> n=A,T,C or G.

<400> 259
gannngagtc acgaaaaggc agtatccctcc tcccccccttt tcgaggactg ccgcacatcttt 60
gttttctgcc cattccagtc accggaaaag atcccaaaga aagaagaaaa gaaacagagg 120
tgcacttcgc ttcatatttc gctcgcttcc ttttctgtct tcacaagtct gcaggattgc 180
ccttgtcctc ttccgagcac atctacgcac gtatgaggtct cgaggtcaaa gccaaaaaaaaa 240
cgcttgcaact cctctttttc ttgcgtgtc tgtgtgtatg tggaattcct tg 292

<210> 260
<211> 582
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(582)
<223> n=A,T,C or G

<400> 260
gcacgagggtt ggggtggtaact gtgtataata actccagatc cttgaccaag tttggagagt 60
cacttatggc catttggaaac caaatgaagg atcaaaggac taattatttt gaataacctct 120
gagtgttttccc caaagcttg agaagagttt cattcagcta taaaatgctc attgtgcaaa 180
tgagtgttttccatgtgtta taattaaagc attgccttta ataataatttt attaccttta 240
gcttgcctttttaatttgag gaaaatccaa acaatttaaa gtaaaacgtg ataaaagacag 300
tttttcngga gananaaggg nagatcgcta tgtttattcc acttaatatc tatatcaaata 360
atttgatca aaagcagact ctcactttaa aaatattttt ctaatggcna gaatcttttn 420
ccttagattga gagtcagagc tcacatagna tnactgctgg taaaatagaca ctttagactat 480
agagctnagc tnaagttcca actanccaaac tgcatatctg aatatgctt ttattnaaag 540
gccagnnctt ttgcctttt nccncctaa tnccttctat tg 582

<210> 261
<211> 783
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(783)
<223> n=A,T,C or G

<400> 261
gcacgaggca aaatacagag ggtatttac catggacagg caacccattt ttccaggaca 60
actcttgca gcagagagct attctcttcc ttttgcctta cactctcaac ctcactcttc 120
gagtgtctgc atccctanttt tccatggcca taagataagg aaccatgagt gttactctag 180
atgaggctgt ttcatgtgg gagctcatcc aggatccaag gtagattcat cagaaggta 240
agtataaggag tggaaaccca aatctctact tttattttga ggccttctct cctcaatttt 300
aaattgtaaa atcaaactta aaactgggtt tctgatggcc agttaaaaaga ctgggtatct 360
gattgccagt taagagatgg tcatttatgc tcaccaccat tctcaagacg caggtgaggt 420
gacangctt ctggggatg ctgancgaat cccccaaatgc cttcaggatt ctggaaatgg 480

tggctctgnt ttaaaactgn tgaactttac aaagaggccta cccgtcatgg ggggactggg 540
aagaaaaacc anangcagt tctggcccan ggttacaccc ccangntac cttgaaggnt 600
tttggacat acctnttncc cccctnttac tgnttcatta gggcntcnnc aacccaantt 660
tccaagttnt ggcccttcna aaanttttt ntttccntt tccanggacc cccctggntt 720
cctggnnccc ccttttata nccaaccttg ccnggnattt ttcncntn aaaggaaat 780
aat 783

<210> 262
<211> 741
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(741)
<223> n=A,T,C or G

<400> 262
tgaaccctan tggcccccgc cccctcgagt cgacggtatac gataagcttg atatcgaatt 60
cggcacgagt gtatattctg ttattatacc ccagattnaa gtgtatattc ttaggcagta 120
gttctgtta acatccttac tacataaaaat ccacttacta tttaagtatt attctaacag 180
gaggtagaat agctgccta aaaaatgtag tgatcgaatg gcagttttc tgctgaatgg 240
aaattactga cacaaaattt ggttttggga gacattttcc tccttgggt tgagtttcc 300
cattcacgga tagggcataa agcttggttt atagttgagg ggtgcaaaag gggaaatagga 360
ttgggaaaat acagtgttcc agcaaaggta tgacaaggta catcttggag aggattccct 420
ttctgtang tggcactgta ngtctgaaa tactgtgtac tttccagaca aaggatagag 480
aaaaagacct tcactgggtg ggggagaaga aaacccttgt tcctagaaaa atcacaaaaa 540
aggcattcct tancctatat tcccagnntt actggngcat ttgcttgatg tgactgacnc 600
ngattatttc cttnactgg naaaaattcc tgccncttg gatatnaang ggggnaccng 660
aaaaatnggg ggcnttgggg aagggaaanaa aaaaattgg agggaccnnaa ctttggaaaa 720
tgggntgctt nangccttaa g 741

<210> 263
<211> 437
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(437)
<223> n=A,T,C or G

<400> 263
ggcacgagag aatgtgttca cagacactat tttatannta tctgatgtgt actgtgtctg 60
gtggatgtga aagccatact tcttaaatct gatttggaaaa gcaaattctga ttatcacagc 120
cataataaa ttggccagc cttcttctt ccctccctcc ttcaacttcc ttcttccttc 180
cgcctcggtc cgaattcgcc acgagcctga cctcaactacc aaaaaaaaaa aaattcaaag 240
tgcctgaggt ttccaggcat tcttagctt atttactac ttcccacctc aaatggcctt 300
agaattcaaa ttctgnanaa aatggattgc catanataat ccaatgaaaa tgggtcatat 360
tttgcattt atagaatcac agtcnacaag ggactaatag aatttagtcac ttangtatcn 420
tttagatttgg gagacnn 437

<210> 264
<211> 706
<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(706)

<223> n=A,T,C or G

<400> 264

gcacgagcac cccaagggtt taggacaaaa tggatgagt gaattcatgg cttgacagac 60
tgaacagaaa aatgaggctc cgtgctccat attcatgtgc atctgcccct catggtgaca 120
tgctaattgg ttggccgggtg cacaagacaa ggaagtgcag gtttcctgtt gtcacacag 180
tgcttcctgt ctgctgtggc aggagccggg aggaaggggag cgagccaaga ggggtgctgc 240
ccaccggaaa cgatggcgcg aggccgcaga gctaaatggg ggccctctcca gggagtgctc 300
tgttcacggc tccatcgctg ttagtaagta tcttgtgatt tcgaaattta aatgaggttg 360
tgtttaacct gcataacatc tggctttaa aatctgactt tatttcctt ttatttcgt 420
gcatcggttc aggcacactt agtggtggct taggtgttga agtcaggtt ccaaacagca 480
cgccctctct ttatttcctag gctgcgtgtt tcattgattc tgaaggtcag atggctgtgt 540
tcaagttctg ttagtatatt ggtgtcagaa atgaaaagat gatgtAACCC ttataactt 600
cttaaaggct catatcatgt cagggaaatta acctgtacga gttatggaca aatgcccattc 660
ctgatgattt tcancatga aaatgaatna aagggganaa gggcca 706

<210> 265

<211> 717

<212> DNA

<213> Homo sapiens

<400> 265

ggcacgagca gcattacggt ttatacacat gtccacaact cagcattgct ttcaaaatag 60
gaacacttta ttagtaaaga ggaagaaatt gcctaaacag actcagtgtc tttccatcaa 120
caatcatctg ccaagccgca ggccttaacca ggaatccca tttccttttgcgttgc 180
ctccaccaac agatacaacc ctgatgccaa atgtttagtgc gttttaggt gttgtgagcc 240
aatgagggca tgccctaggggc caaaggctgc cctttggaaat gaggggcaagg tcgttagactc 300
catcaaacaa caaatgcattc ctccctccaa atcaaatgtc caacacatgc agccttcgt 360
atgcccattt cccctttact cattttcatg gctgaaaatc atcaggatgg gcatttgc 420
ataactccctt caggttaatt tcctgacatg atatgagctt ttaagaagtt ataaagggtt 480
acatcatctt ttcatattctg acaccaatat actaacagaa cttaaacaca gccatctgac 540
cttcagaatc aatgaaacac gcagcctgag aataaagaga gggcgtgtt ttggtaacc 600
tgacttcaac acctaagcca ccactaagtg tgcctgagcc gatgcacaga aataaaagga 660
aaataaagtc agattttaaa aagccagatg ttatgcaggg taaacacaac ctcattt 717

<210> 266

<211> 362

<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(362)

<223> n=A,T,C or G

<400> 266

ggcacgaggt tagatttaac ttccacagat gactcagcag aggataacta ctaatcagag 60
tacaacatca aaactgtaac cagtataatc actggattat gagcaactca aaatagctcc 120
agtttccaaa gggccataaa ctgcacatcat cagtactatg tgcaattaac acataattta 180
ttatgaaaat gtggacatgc caggttaagta aggggattta gtttgacttt ttataataact 240
ttaaatttga aatgccattt ctgtggattt gatgacatct tccaggtgtt ntaatnctgg 300

gntacctnct gatanatcct gananaaaaga ggtancacca gcgtctatca nacctcaata 360
ca 362

<210> 267
<211> 692
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(692)
<223> n=A,T,C or G

<400> 267
ggcacaggt tagatttaac ttccacagat gactcagcag aggataacta ctaatcagag 60
tacaacatca aaactgtAAC cagtataatc actggattat gagcaactca aaatagctcc 120
agtttccaaa gggccataaac tggccctttt aanactttnn gcaattaaca cataatttat 180
tatgaaaatg tggacatgcc aggttaagtaa ggggatttag gtgactttt tataataactt 240
taaatttcaa atgcatttc tgtggatttg atgacatctt ccaggtgctt taatttggtt 300
tacccctga tagatcctga cagaaagagg nacaccagc gtctatcaaa cctcaataca 360
gngtgtgaaa cacangagag cctgcttttgc tcnacacggg gaaacacatt gttatcacaa 420
cacacaaaag gcaanctncc aatgggnan ncttacctgn cctctcatat tggggcaan 480
gaaaangggg ccccccanatg gctgagtana tcccaaaaaa ccnccactan tggtcagnnt 540
gcttccccan acagccagat gactgaattt agcccaagct gcagtctcaa aaccagctt 600
ctgacaatca gtaacaagaa catactggtc tggcagtg agctcaagtg ttgggtgttc 660
agtcaaaaanc catggatgcc aatcatctcc ca 692

<210> 268
<211> 605
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(605)
<223> n=A,T,C or G

<400> 268
cgtgccaaat tcggcacgag ngcacatatac agtactatgt gcaattaaca cataatttat 60
tatgaaaatg tggacatgcc aggttaagtaa ggggatttan gtgactttt tataataactt 120
taaatttcaa atgcatttc tgtggatttg atgacatctt ccaggtgctt taatttggtt 180
tacccctga tagatcctga cagaaagagg tagcaccagc gtctatcaaa cctcaataca 240
gttgtaaaac acagagagcc tgcttccta cacatggaga aacattgtta tcacaagaca 300
cagaaggcaa acttccaatc tggcataactt ncctgtcctc tcataatttg ggcaatgaga 360
atggtgacc agatggctt antagatgcc aaagaacacc canactgggc agcatgctt 420
cccagacagc cngaaagactg aaatttantic ccagctgcag ncttaaaacc ttttttgac 480
nttccgtaac cagaccatac tttttttct gatgcttttc ttaacttcat cttttccaat 540
taaattcatt agtnnaaccc taaangggc cggtttccg aaaaatttcc ntntnttt 600
ccccn 605

<210> 269
<211> 535
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(535)
<223> n=A,T,C or G

<400> 269

gcacgagggc aaccccagg gtgggtctc tggatgaac tggagacct gagcttcac 60
agttcccttg taaaatttagg gaggcatggc ccacaagatt ccaagctcc ttcttatcca 120
aacttgcatt ttttagattt catgtccag ttcacacgg ttatggctg aatctcatgc 180
actanaaaaaa ggtaatataa aaganaaaaa tanaangatn ttcaagttag tataaanacc 240
tttaatctca ntctttctag ttcaaagaga cggacaatg agagatgctg gttcatanag 300
ctgntanatt taacttccac agatgactca ncagaggata actactaattc anagtacaac 360
atcaaaaactg taaccagtat aatcactgga ttatgagcaa ctcaaaaatag ctccagttc 420
caaagggcca taaactgcca tatcaantac tatgtgccat taaccataa ttattatga 480
aatgtggac atgcccangtn agtaagggga tttagggta cttttatna tactt 535

<210> 270

<211> 803
<212> DNA
<213> Homo sapiens

<220>

<221> misc_feature
<222> (1)...(803)
<223> n=A,T,C or G

<400> 270

gcacgagggc aaccccaggg tggttctct gggatgaacc tggagacctg agcttcac 60
gcttccttgg taaaatttaggg aggcatggac cacaagattt ccaagctct ttcttatccaa 120
acttgcattt ttttagattt atgtccagt tcatcacgg ttagggctga atctcatgca 180
ctagaaaaaa gtaatataaa agaaaaaaat aaaaagatatt tcaagttagt ataaagacct 240
ttaatctcag tctttcttagt tcaaagagac ggaacaatga gagatgctgg ttcataagac 300
tgttagattt aacttccaca gatgactcag cagaggataa ctactaatca gagtacaaca 360
tcaaaaactgt aaccagtata atcactggat tatgagcaac tcaaaatagc tccagttcc 420
aaagggccat aaactgcaca tatcagtact atgtcaatt aacacataat ttattatgaa 480
aatgtggaca tgccaggtaa gtaagggat tttaggttgc tttttataat actttaaatt 540
tgaaatgcca ttctgtgga ttggatgaca tcttccaggt gcttaattt ggtttacctc 600
ctgatagatc ctgacagaaa gagtagcac cagcgcttat caaacctcaa tacagttgta 660
aaacacagag agcctgnntt gcctacncat ggagaacatt gttatcacaa gacacagaag 720
ggaacttcca tctggctact tacctggctt tattttttggg gcaatganaa tngggggacc 780
aatggntgan tanatgccaa aaa 803

<210> 271

<211> 836
<212> DNA
<213> Homo sapiens

<220>

<221> misc_feature
<222> (1)...(836)
<223> n=A,T,C or G

<400> 271

gcacgagggc aaccccaggg tggttctct gggatgaacc tggagacctg agcttcac 60
gcttccttgg taaaatttaggg aggcatggac cacaagattt ccaagctct ttcttatccaa 120
acttgcattt ttttagattt atgtccagt tcatcacgg ttagggctga atctcatgca 180

ctagaaaaaag gtaatataaa agaaaaaaaaat aaaaagatat tcaagtgagt ataaagacct 240
ttaatctcag tctttcttagt tcaaagagac ggaacaatga gagatgctgg ttcatalogc 300
tgtagattt aactccaca gatgactcag cagaggataa ctactaatca gagtacaaca 360
tcaaaactgt aaccagtata atcactggat tatgagcaac tcaaaatagc tccagttcc 420
aaagggccat aaactgcaca tatcagtact atgtgcaatt aacacataat ttattatgaa 480
aatgtggaca tgccaggtaa gtaaggggat ttaggttgac ttttataat actttaaatt 540
tgaatgcca tttctgtgga ttggatgaca tcttcaggt gctttaattt ggtttacctc 600
ctgatagatc ctgacagaaa gangtagcac cagcgctat caaacctcaa tacagttgta 660
aaacacagag agcctgcctt gnctacacat ggagaaacat tgtatcacaa gacacagnaa 720
ggcaacttcc atctggata ctacctgtct ctctatttgg ggcatganat gggacaatg 780
ntgananatg caanacacca atngagctg nttccnacag cnatatgatt ntccat 836

<210> 272

<211> 203

<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(203)

<223> n=A,T,C or G

<400> 272

ggagaattgg gcccgtcang ggtgcattct gcatcacctg anttcnaaat ctnagtcaat 60
cnncgtacta atantatcaa catnattta acctgatctc cactgcttng tnatttcnn 120
ttcactgncc ctntcactng aacntctntt cacacagcca ccccccatta tctggntggc 180
accccncca aatnccncc naa 203

<210> 273

<211> 594

<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(594)

<223> n=A,T,C or G

<400> 273

attcgggccn ctggatncgt gctcgagcgg ccggccgtgt gatggatatac tgcanattc 60
ggcttctgga gagagcttn tttttagatgg ttgcangtac tctcgatgga gttgggtgggt 120
gtggttatct ctctctgggt gtctttctgt ataaanttct tgcnctgact ncctanctn 180
cctccccctg gtccttcct tagntaaca nctggtaatc cctntcttct ttgctctcct 240
tncttctcct gancgattc ctctntttgt ccactctcag gnanaaccct gntggtcagt 300
gttcatgact tcnngaagnt cgaccgcna aatagggncn cacggatnat gttgaancng 360
ggaagggagn gtccaanntc tctgttccan aggctnagcc tagaganaat gatgggagan 420
ggtttactga gatcatngnn tcttctcgaa gatatnntt agggtggtcc cccataagng 480
aatttctcan cttcaaattct tctaatacat tactgaacan ctgnccatttgc ttacgccaca 540
nattgnaattt ctccatntct ttttagaaac nattncaagg tcatttattt ccct 594

<210> 274

<211> 229

<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature
<222> (1)...(229)
<223> n=A,T,C or G

<400> 274
ctactcaactg tccggccatt tggncctctg natgcatnct caagcagcnc gccantatga 60
tnnatatctg cacanttcag cttctngaga aaactatgtt taaaacagtgt gentanactt 120
anaatanaaaa tcgagtaagg tntagatnan tctctaaccga tngaattatt ntacanaggg 180
gtanncgatn accaggagta nctaganttg ancancancc taggtcnga 229

<210> 275
<211> 651
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(651)
<223> n=A,T,C or G

<400> 275
atatctgn tg aatacggntt cctgnaaaaa ggtntnattt agatggttga gtccgactca 60
gcgatgcgac ttgggtgggtg tggtcantct cttatgggtt agattgttca tgatcatcg 120
ccctgagatg cctggactnn cctcaccgga gatcctagac ggtgntanc cctgagagtc 180
tctctcntcc tgctctccta acttctccta atgatccctc cnattgtcta ctgtccnatt 240
gaacccttct tgcttatgta tncaatcntt nacgggttcc ctgctnantt tttganacga 300
ngctcataat ggacngggga aggatagtn gaataatntc ctgtataacc acgcnacnt 360
ctacnctntg atctgacacg gtatactgat ttgtgctgtt cncttcacca ttccannttc 420
taccccccgc teatatgctc tgtangctac accctctgtg actgctttct cagttacgtg 480
caacaaggtn ttcatatctn gaactcttac accattctag anggatcncc cctcgganaa 540
annttgaan aacaaggcaag ancanaatnc ctctctngtg ntacacnanc cggttncgt 600
atcctcgtn aaggaattcc ccgccttcct gggcttaan tctcctaaac t 651

<210> 276
<211> 392
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(392)
<223> n=A,T,C or G

<400> 276
accccccgg aattacgntg gcncatntaa aagtnccatca ngcctccang caacntatcn 60
tttcattacc acccacactc ctgttnnggg anggangtgg naatccttca ccatnctaatt 120
gtatgtggtg ctctcatgcn ggtacgtata atctannctg cccctnaaat cggtatgttc 180
tgtaatcnnc agtcacnaaaa ccacangan caactgaaac angatttggc taacagccaa 240
tgtctggcc ctcncnaatc cctnnaatat ctcctacacc tgttagtanna atnaactacn 300
ctacnctatt nnacacacgn tttaggtgt annaccaagc ccttatttgg tagaaatcggtt 360
tntatngtat naaatgcca aagntgcgg aa 392

<210> 277
<211> 212
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(212)
<223> n=A,T,C or G

<400> 277
ggtttgcggg natgaanttt gnaanaatna acttagtnga taacccaccc accaatncct 60
nctnagtatt tgncaacctn aaaactacag ctctctccag atagactntn ccttnctgat 120
ttcaactctc cttggactgg tcagcctgaa gggtggtaat gactcaccaa cgctactaat 180
ncctnttna ctgtgccttn atttttcgc ct 212

<210> 278
<211> 269
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(269)
<223> n=A,T,C or G

<400> 278
nnntccatcc taataccact cactatcggg ctcgaancgg ccgcccgggc acgtntcttn 60
tgngacagga tctgaatnaa gggtggtttg taacttnact naaaattctg aaatgatcct 120
gcatcagaca gggttctccg tntanaatan agttccctg ttagttatcn agcctgggca 180
ggggangana gattcgagga cntntgaaat gaagggnatta tttaggatgg gtgactcatt 240
ccnaccnttc ncgctnacca gnccganga 269

<210> 279
<211> 266
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(266)
<223> n=A,T,C or G

<400> 279
gttggtgant cngttggng tcttccttgt gntnggtgtt tggtgtgttg nnttggtn 60
gggtngtnntt tntggagaga gttgttagttc gtgagggttg cagtgtactt actatggagc 120
ctaaggangt gngctaactt anantgatna ctttgctcat actgccctgc cctnaatgcc 180
nngcttgcct caccctggtg ccnaaccnna tcgaacacct aacagtctag taggcttctt 240
gctntancag actnctctg aggatc 266

<210> 280
<211> 317
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(317)
<223> n=A,T,C or G

<400> 280
acactgtnag gtgtntggaa ntgnntgtagg catagncttt ntggcacaga gttggagccg 60
tgaggcatag cngtactta ctatggagcc taaggangga gctaacttat antnatnact 120
ttgctcatac tgccctgcctc tnaatgccta ngcttgccctc accctgntgc cttacnnnat 180
cgaacaccta cgccgtctat aggcttcttg ctctatcagg actncttcc nagcttcntc 240
gcctcanttg actcaactgtg ctcggcggtt ctactngat ccagncgctc atnaacctna 300
cttnggacgc aggtcat 317

<210> 281
<211> 174
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(174)
<223> n=A,T,C or G

<400> 281
gnggtcatat tatacatcta aggcatggcc aactccacgc cattatnaat tccatcgta 60
tgtccgcagt cactacttat aaccttagatt aatagtgcct ggccccggac ngtctgtca 120
atctnccgcc ataccaattn cgatccncan accncgatna cactcctcct tact 174

<210> 282
<211> 169
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(169)
<223> n=A,T,C or G

<400> 282
atcgcaagtt gtacgatcgt catataacgc gcatgtgcgg atcgcttcag cgccgcccga 60
ctgtcagaag gangagatct ttttatcac ttgtttgtt gactatanat aanancgact 120
acagcattga tgtgtgtcct caaganttg ctgggtctga naaagctga 169

<210> 283
<211> 157
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(157)
<223> n=A,T,C or G

<400> 283
ggntntctaa gatcgcaagg gtacgatcgt catatnacgc gcatgtgcgn atcgcttcac 60
gtcgccnggc tgtccaggan atgcatntca acataatgtg cactctatat ggttattgat 120
taatacgagn tangagcana tatcngatac aacacaa 157

<210> 284
<211> 133
<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(133)

<223> n=A,T,C or G

<400> 284

ggngtgggt nagatacgca ngctgggacg aatcgntca tagtacggcg catgtgttga 60
tcaattctga aaatccatcc cggcgcgcctc ancatgact anaggcaat cgccatatatg 120
antcgattt caa 133

<210> 285

<211> 194

<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(194)

<223> n=A,T,C or G

<400> 285

ntntgngtga tgataccaa gctggntacc nactngantc caattaccgg ctcantntgc 60
tnaaaaacngc ttcgatngnc tcctggcatg tacttggaaac aggnatanata tctaataagnn 120
tacngtgnnn tttcnatca tacagnttt atatncact ncctnccatt cttttctant 180
ctctctctcc ntat 194

<210> 286

<211> 134

<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(134)

<223> n=A,T,C or G

<400> 286

gagggnnat gataccaagc tggtaganc ccgtcaactat nacggcccag tgtgtggatc 60
cgctanctgg tcnccgatg tctacncaca cngaaactgc ctctcgnaa gatctcctct 120
cctctccnaa gaga 134

<210> 287

<211> 119

<212> DNA

<213> Homo sapiens

<220>

<221> misc_feature

<222> (1)...(119)

<223> n=A,T,C or G

<400> 287

tnggttat ccagttgtac actggncata tacgcgcatt atgatcgaaa cacgccccga 60
gtacggcattc attacganat ggnctcattc gtttaccttt ntcgctggac acaagcgta 119

<210> 288

<211> 170
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(170)
<223> n=A,T,C or G

<400> 288
gggntgagat acncaagttg gtacgagtcg gatcatatna cggncgccat tttcttgaat 60
ccgcttacgt ggtcccgccg aagtactttt tcattgccttg caaaatngcg ttactgcact 120
ancttgccta acctatgagt ggggtcttcc ataccncntc tntcatggaa 170

<210> 289
<211> 126
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(126)
<223> n=A,T,C or G

<400> 289
ggccaattgg ggctctana tgcntgctcg aacgggcgcc aatttnatgg atatctccaa 60
aattcggctt accntggtcg cggncnaagt acttaactca atccatctnt cactcaggat 120
naatgc 126

<210> 290
<211> 126
<212> DNA
<213> Homo sapiens

<220>
<221> misc_feature
<222> (1)...(126)
<223> n=A,T,C or G

<400> 290
ggccaattgg ggctctana tgcntgctcg aacgggcgcc aatttnatgg atatctccaa 60
aattcggctt accntggtcg cggncnaagt acttaactca atccatctnt cactcaggat 120
naatgc 126

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